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(54) Title: INDUCING CELLULAR IMMUNE RESPONSES TO HEPATITIS C VIRUS USING PEPTIDE AND NUCLEIC ACID COMPOSITIONS

(57) Abstract: This invention uses our knowledge of the mechanisms by which antigen is recognized by T cells to identify and prepare HCV epitopes, and to develop epitope-based vaccines directed towards HCV. More specifically, this application communicates our discovery of pharmaceutical compositions and methods of use in the prevention and treatment of HCV infection.

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**INDUCING CELLULAR IMMUNE RESPONSES TO HEPATITIS C VIRUS  
USING PEPTIDE AND NUCLEIC ACID COMPOSITIONS**

**FEDERALLY SPONSORED RESEARCH AND DEVELOPMENT**

10 This invention was funded, in part, by the United States government under grants with the National Institutes of Health. The U.S. government has certain rights in this invention.

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VII. Abstract

## I. BACKGROUND OF THE INVENTION

Hepatitis C virus (HCV) infection is a global human health problem with approximately 150,000 new reported cases each year in the U.S. alone. HCV is a single stranded RNA virus, and is the etiological agent identified in most cases of non-A, non-B post-transfusion and post-transplant hepatitis, and is a common cause of acute sporadic hepatitis (Choo *et al.*, *Science* 244:359, 1989; Kuo *et al.*, *Science* 244:362, 1989; and Alter *et al.*, in: *Current Perspective in Hepatology*, p. 83, 1989). It is estimated that more than 50% of patients infected with HCV become chronically infected and, of those, 20% develop cirrhosis of the liver within 20 years (Davis *et al.*, *New Engl. J. Med.* 321:1501,

1989; Alter *et al.*, in: *Current Perspective in Hepatology*, p. 83, 1989; Alter *et al.*, *New Engl. J. Med.* 327:1899, 1992; and Dienstag, J. L. *Gastroenterology* 85:430, 1983). Moreover, the only therapy available for treatment of HCV infection is interferon- $\alpha$ . Most patients are unresponsive, however, and among the responders, there is a high 5 recurrence rate within 6-12 months of cessation of treatment (Liang *et al.*, *J. Med. Virol.* 40:69, 1993). Ribaviron, a guanosine analog with a broad spectrum activity against many RNA and DNA viruses, has been shown in clinical trials to be effective against chronic HCV infection when used in combination with interferon-  $\alpha$  (see, e.g., Poynard *et al.*, *Lancet* 352:1426-1432, 1998; Reichard *et al.*, *Lancet* 351:83-87, 1998) However, the 10 response rate is still well below 50%.

Virus-specific, human leukocyte antigen (HLA) class I-restricted cytotoxic T lymphocytes (CTL) are known to play a major role in the prevention and clearance of virus infections *in vivo* (Oldstone *et al.*, *Nature* 321:239, 1989; Jamieson *et al.*, *J. Virol.* 61:3930, 1987; Yap *et al.*, *Nature* 273:238, 1978; Lukacher *et al.*, *J. Exp. Med.* 160:814, 15 1994; McMichael *et al.*, *N. Engl. J. Med.* 309:13, 1983; Sethi *et al.*, *J. Gen. Virol.* 64:443, 1983; Watari *et al.*, *J. Exp. Med.* 165:459, 1987; Yasukawa *et al.*, *J. Immunol.* 143:2051, 1989; Tigges *et al.*, *J. Virol.* 66:1622, 1993; Reddenhase *et al.*, *J. Virol.* 55:263, 1985; Quinnan *et al.*, *N. Engl. J. Med.* 307:6, 1982). HLA class I molecules are expressed on the surface of almost all nucleated cells. Following intracellular processing of antigens, 20 epitopes from the antigens are presented as a complex with the HLA class I molecules on the surface of such cells. CTL recognize the peptide-HLA class I complex, which then results in the destruction of the cell bearing the HLA-peptide complex directly by the CTL and/or via the activation of non-destructive mechanisms e.g., the production of interferon, that inhibit viral replication.

25 In view of the heterogeneous immune response observed with HCV infection, induction of a multi-specific cellular immune response directed simultaneously against multiple HCV epitopes appears to be important for the development of an efficacious vaccine against HCV. There is a need, however, to establish vaccine embodiments that elicit immune responses that correspond to responses seen in patients that clear HCV 30 infection.

The information provided in this section is intended to disclose the presently understood state of the art as of the filing date of the present application. Information is included in this section which was generated subsequent to the priority date of this

application. Accordingly, information in this section is not intended, in any way, to delineate the priority date for the invention.

## II. SUMMARY OF THE INVENTION

5 This invention applies our knowledge of the mechanisms by which antigen is recognized by T cells, for example, to develop epitope-based vaccines directed towards HCV. More specifically, this application communicates our discovery of specific epitope pharmaceutical compositions and methods of use in the prevention and treatment of HCV infection.

10 Upon development of appropriate technology, the use of epitope-based vaccines has several advantages over current vaccines, particularly when compared to the use of whole antigens in vaccine compositions. There is evidence that the immune response to whole antigens is directed largely toward variable regions of the antigen, allowing for immune escape due to mutations. The epitopes for inclusion in an epitope-based vaccine 15 are selected from conserved regions of viral or tumor-associated antigens, which thereby reduces the likelihood of escape mutants. Furthermore, immunosuppressive epitopes that may be present in whole antigens can be avoided with the use of epitope-based vaccines.

20 An additional advantage of an epitope-based vaccine approach is the ability to combine selected epitopes (CTL and HTL), and further, to modify the composition of the epitopes, achieving, for example, enhanced immunogenicity. Accordingly, the immune response can be modulated, as appropriate, for the target disease. Similar engineering of the response is not possible with traditional approaches.

25 Another major benefit of epitope-based immune-stimulating vaccines is their safety. The possible pathological side effects caused by infectious agents or whole protein antigens, which might have their own intrinsic biological activity, is eliminated.

An epitope-based vaccine also provides the ability to direct and focus an immune response to multiple selected antigens from the same pathogen. Thus, patient-by-patient variability in the immune response to a particular pathogen may be alleviated by inclusion of epitopes from multiple antigens from that pathogen in a vaccine composition. A 30 "pathogen" may be an infectious agent or a tumor associated molecule.

One of the most formidable obstacles to the development of broadly efficacious epitope-based immunotherapeutics, however, has been the extreme polymorphism of HLA molecules. To date, effective non-genetically biased coverage of a population has been a task of considerable complexity; such coverage has required that epitopes be used

that are specific for HLA molecules corresponding to each individual HLA allele, therefore, impractically large numbers of epitopes would have to be used in order to cover ethnically diverse populations. Thus, there has existed a need for peptide epitopes that are bound by multiple HLA antigen molecules for use in epitope-based vaccines. The 5 greater the number of HLA antigen molecules bound, the greater the breadth of population coverage by the vaccine.

Furthermore, as described herein in greater detail, a need has existed to modulate peptide binding properties, for example, so that peptides that are able to bind to multiple HLA antigens do so with an affinity that will stimulate an immune response. 10 Identification of epitopes restricted by more than one HLA allele at an affinity that correlates with immunogenicity is important to provide thorough population coverage, and to allow the elicitation of responses of sufficient vigor to prevent or clear an infection in a diverse segment of the population. Such a response can also target a broad array of epitopes. The technology disclosed herein provides for such favored immune responses.

15 In a preferred embodiment, epitopes for inclusion in vaccine compositions of the invention are selected by a process whereby protein sequences of known antigens are evaluated for the presence of motif or supermotif-bearing epitopes. Peptides corresponding to a motif- or supermotif-bearing epitope are then synthesized and tested for the ability to bind to the HLA molecule that recognizes the selected motif. Those 20 peptides that bind at an intermediate or high affinity *i.e.*, an  $IC_{50}$  (or a  $K_D$  value) of 500 nM or less for HLA class I molecules or 1000 nM or less for HLA class II molecules, are further evaluated for their ability to induce a CTL or HTL response. Immunogenic peptide epitopes are selected for inclusion in vaccine compositions.

25 Supermotif-bearing peptides may additionally be tested for the ability to bind to multiple alleles within the HLA supertype family. Moreover, peptide epitopes may be analogued to modify binding affinity and/or the ability to bind to multiple alleles within an HLA supertype.

The invention also includes an embodiment comprising a method for monitoring or evaluating an immune response to HCV in a patient having a known HLA-type, the 30 method comprising incubating a T lymphocyte sample from the patient with a peptide composition comprising an HCV epitope consisting essentially of an amino acid sequence described in Tables VII to Table XX or Table XXII which binds the product of at least one HLA allele present in said patient, and detecting for the presence of a T lymphocyte

that binds to the peptide. A CTL peptide epitope may, for example, comprise a tetrameric complex.

An alternative modality for defining the peptide epitopes in accordance with the invention is to recite the physical properties, such as length; primary structure; or charge, which are correlated with binding to a particular allele-specific HLA molecule or group of allele-specific HLA molecules. A further modality for defining peptide epitopes is to recite the physical properties of an HLA binding pocket, or properties shared by several allele-specific HLA binding pockets (e.g. pocket configuration and charge distribution) and reciting that the peptide epitope fits and binds to said pocket or pockets.

10 As will be apparent from the discussion below, other methods and embodiments are also contemplated. Further, novel synthetic peptides produced by any of the methods described herein are also part of the invention.

### III. BRIEF DESCRIPTION OF THE FIGURES

15 Figure 1: Figure 1 provides a graph of total frequency of genotypes as a function of the number of HCV candidate epitopes bound by HLA-A and B molecules, in an average population.

20 Figure 2: Figure 2 illustrates the position of peptide epitopes in an experimental model minigene construct.

### IV. DETAILED DESCRIPTION OF THE INVENTION

The peptide epitopes and corresponding nucleic acid compositions of the present invention are useful for stimulating an immune response to HCV by stimulating the production of CTL or HTL responses. The peptide epitopes, which are derived directly or indirectly from native HCV amino acid sequences, are able to bind to HLA molecules and stimulate an immune response to HCV. The complete polyprotein sequence from HCV and its variants can be obtained from Genbank. Peptide epitopes and analogs thereof can also be readily determined from sequence information that may subsequently be discovered for heretofore unknown variants of HCV, as will be clear from the disclosure provided below.

The peptide epitopes of the invention have been identified in a number of ways, as will be discussed below. Also discussed in greater detail is that analog peptides have been derived and the binding activity for HLA molecules modulated by modifying specific amino acid residues to create peptide analogs exhibiting altered immunogenicity.

Further, the present invention provides compositions and combinations of compositions that enable epitope-based vaccines that are capable of interacting with HLA molecules encoded by various genetic alleles to provide broader population coverage than prior vaccines.

5

#### IV.A. Definitions

The invention can be better understood with reference to the following definitions, which are listed alphabetically:

A "computer" or "computer system" generally includes: a processor; at least one information storage/retrieval apparatus such as, for example, a hard drive, a disk drive or a tape drive; at least one input apparatus such as, for example, a keyboard, a mouse, a touch screen, or a microphone; and display structure. Additionally, the computer may include a communication channel in communication with a network. Such a computer may include more or less than what is listed above.

15 "Cross-reactive binding" indicates that a peptide is bound by more than one HLA molecule; a synonym is degenerate binding.

A "cryptic epitope" elicits a response by immunization with an isolated peptide, but the response is not cross-reactive *in vitro* when intact whole protein which comprises the epitope is used as an antigen.

20 A "dominant epitope" is an epitope that induces an immune response upon immunization with a whole native antigen (see, e.g., Sercarz, *et al.*, *Annu. Rev. Immunol.* 11:729-766, 1993). Such a response is cross-reactive *in vitro* with an isolated peptide epitope.

25 With regard to a particular amino acid sequence, an "epitope" is a set of amino acid residues which is involved in recognition by a particular immunoglobulin, or in the context of T cells, those residues necessary for recognition by T cell receptor proteins and/or Major Histocompatibility Complex (MHC) receptors. In an immune system setting, *in vivo* or *in vitro*, an epitope is the collective features of a molecule, such as primary, secondary and tertiary peptide structure, and charge, that together form a site 30 recognized by an immunoglobulin, T cell receptor or HLA molecule. Throughout this disclosure epitope and peptide are often used interchangeably.

It is to be appreciated that protein or peptide molecules that comprise an epitope of the invention as well as additional amino acid(s) are still within the bounds of the invention. In certain embodiments, there is a limitation on the length of a peptide of the

invention which is not otherwise a construct. An embodiment that is length-limited occurs when the protein/peptide comprising an epitope of the invention comprises a region (i.e., a contiguous series of amino acids) having 100% identity with a native sequence. In order to avoid the definition of epitope from reading, e.g., on whole natural molecules, there is a limitation on the length of any region that has 100% identity with a native peptide sequence. Thus, for a peptide comprising an epitope of the invention and a region with 100% identity with a native peptide sequence (and is not otherwise a construct), the region with 100% identity to a native sequence generally has a length of: less than or equal to 600 amino acids, often less than or equal to 500 amino acids, often less than or equal to 400 amino acids, often less than or equal to 250 amino acids, often less than or equal to 100 amino acids, often less than or equal to 85 amino acids, often less than or equal to 75 amino acids, often less than or equal to 65 amino acids, and often less than or equal to 50 amino acids. In certain embodiments, an "epitope" of the invention is comprised by a peptide having a region with less than 51 amino acids that has 100% identity to a native peptide sequence, in any increment of (49, 48, 47, 46, 45, 44, 43, 42, 41, 40, 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5) down to 5 amino acids.

Accordingly, peptide or protein sequences longer than 600 amino acids are within the scope of the invention, so long as they do not comprise any contiguous sequence of more than 600 amino acids that have 100% identity with a native peptide sequence, if they are not otherwise a construct. For any peptide that has five contiguous residues or less that correspond to a native sequence, there is no limitation on the maximal length of that peptide in order to fall within the scope of the invention. It is presently preferred that a CTL epitope be less than 600 residues long in any increment down to eight amino acid residues.

"Human Leukocyte Antigen" or "HLA" is a human class I or class II Major Histocompatibility Complex (MHC) protein (*see, e.g.*, Stites, *et al.*, IMMUNOLOGY, 8<sup>TH</sup> ED., Lange Publishing, Los Altos, CA (1994)).

An "HLA supertype or family", as used herein, describes sets of HLA molecules grouped on the basis of shared peptide-binding specificities. HLA class I molecules that share somewhat similar binding affinity for peptides bearing certain amino acid motifs are grouped into HLA supertypes. The terms HLA superfamily, HLA supertype family, HLA family, and HLA xx-like supertype molecules (where xx denotes a particular HLA type), are synonyms.

Throughout this disclosure, results are expressed in terms of "IC<sub>50</sub>'s." IC<sub>50</sub> is the concentration of peptide in a binding assay at which 50% inhibition of binding of a reference peptide is observed. Given the conditions in which the assays are run (*i.e.*, limiting HLA proteins and labeled peptide concentrations), these values approximate K<sub>D</sub> values. Assays for determining binding are described in detail, *e.g.*, in PCT publications WO 94/20127 and WO 94/03205. It should be noted that IC<sub>50</sub> values can change, often dramatically, if the assay conditions are varied, and depending on the particular reagents used (*e.g.*, HLA preparation, *etc.*). For example, excessive concentrations of HLA molecules will increase the apparent measured IC<sub>50</sub> of a given ligand.

10 Alternatively, binding is expressed relative to a reference peptide. Although as a particular assay becomes more, or less, sensitive, the IC<sub>50</sub>'s of the peptides tested may change somewhat, the binding relative to the reference peptide will not significantly change. For example, in an assay run under conditions such that the IC<sub>50</sub> of the reference peptide increases 10-fold, the IC<sub>50</sub> values of the test peptides will also shift approximately 15 10-fold. Therefore, to avoid ambiguities, the assessment of whether a peptide is a good, intermediate, weak, or negative binder is generally based on its IC<sub>50</sub>, relative to the IC<sub>50</sub> of a standard peptide.

Binding may also be determined using other assay systems including those using: live cells (*e.g.*, Ceppellini *et al.*, *Nature* 339:392, 1989; Christnick *et al.*, *Nature* 352:67, 20 1991; Busch *et al.*, *Int. Immunol.* 2:443, 1999; Hill *et al.*, *J. Immunol.* 147:189, 1991; del Guercio *et al.*, *J. Immunol.* 154:685, 1995), cell free systems using detergent lysates (*e.g.*, Cerundolo *et al.*, *J. Immunol.* 21:2069, 1991), immobilized purified MHC (*e.g.*, Hill *et al.*, *J. Immunol.* 152, 2890, 1994; Marshall *et al.*, *J. Immunol.* 152:4946, 1994), ELISA systems (*e.g.*, Reay *et al.*, *EMBO J.* 11:2829, 1992), surface plasmon resonance (*e.g.*, 25 Khilko *et al.*, *J. Biol. Chem.* 268:15425, 1993); high flux soluble phase assays (Hammer *et al.*, *J. Exp. Med.* 180:2353, 1994), and measurement of class I MHC stabilization or assembly (*e.g.*, Ljunggren *et al.*, *Nature* 346:476, 1990; Schumacher *et al.*, *Cell* 62:563, 1990; Townsend *et al.*, *Cell* 62:285, 1990; Parker *et al.*, *J. Immunol.* 149:1896, 1992).

As used herein, "high affinity" with respect to HLA class I molecules is defined as 30 binding with an IC<sub>50</sub>, or K<sub>D</sub> value, of 50 nM or less; "intermediate affinity" is binding with an IC<sub>50</sub> or K<sub>D</sub> value of between about 50 and about 500 nM. "High affinity" with respect to binding to HLA class II molecules is defined as binding with an IC<sub>50</sub> or K<sub>D</sub> value of 100 nM or less; "intermediate affinity" is binding with an IC<sub>50</sub> or K<sub>D</sub> value of between about 100 and about 1000 nM.

The terms "identical" or percent "identity," in the context of two or more peptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues that are the same, when compared and aligned for maximum correspondence over a comparison window, as measured using a sequence comparison algorithm or by manual alignment and visual inspection.

An "immunogenic peptide" or "peptide epitope" is a peptide that comprises an allele-specific motif or supermotif such that the peptide will bind an HLA molecule and induce a CTL and/or HTL response. Thus, immunogenic peptides of the invention are capable of binding to an appropriate HLA molecule and thereafter inducing an HLA-restricted cytotoxic or helper T cell response to the antigen from which the immunogenic peptide is derived.

The phrases "isolated" or "biologically pure" refer to material which is substantially or essentially free from components which normally accompany the material as it is found in its native state. Thus, isolated peptides in accordance with the invention preferably do not contain materials normally associated with the peptides in their *in situ* environment. An "isolated" epitope refers to an epitope that does not include the whole sequence of the antigen or polypeptide from which the epitope was derived. Typically the "isolated" epitope does not have attached thereto additional amino acids that result in a sequence that has 100% identity with a native sequence. The native sequence can be a sequence such as a tumor-associated antigen from which the epitope is derived.

"Major Histocompatibility Complex" or "MHC" is a cluster of genes that plays a role in control of the cellular interactions responsible for physiologic immune responses. In humans, the MHC complex is also known as the HLA complex. For a detailed description of the MHC and HLA complexes, see, Paul, FUNDAMENTAL IMMUNOLOGY, 25 3<sup>RD</sup> ED., Raven Press, New York, 1993.

The term "motif" refers to the pattern of residues in a peptide of defined length, usually a peptide of from about 8 to about 13 amino acids for a class I HLA motif and from about 6 to about 25 amino acids for a class II HLA motif, which is recognized by a particular HLA molecule. Peptide motifs are typically different for each protein encoded by each human HLA allele and differ in the pattern of the primary and secondary anchor residues.

A "negative binding residue" is an amino acid which, if present at certain positions (typically not primary anchor positions) in a peptide epitope, results in decreased binding affinity of the peptide for the peptide's corresponding HLA molecule.

A "non-native" sequence or "construct" refers to a sequence that is not found in in nature ("non-naturally occurring"). Such sequences include, *e.g.*, peptides that are lipidated or otherwise modified and polyepitopic compositions that contain epitopes that are non contiguous in a native protein sequence.

5 The term "peptide" is used interchangeably with "oligopeptide" in the present specification to designate a series of residues, typically L-amino acids, connected one to the other, typically by peptide bonds between the  $\alpha$ -amino and carboxyl groups of adjacent amino acids. The preferred CTL-inducing peptides of the invention are 13 residues or less in length and usually consist of between about 8 and about 11 residues, 10 preferably 9 or 10 residues. The preferred HTL-inducing oligopeptides are less than about 50 residues in length and usually consist of between about 6 and about 30 residues, more usually between about 12 and 25, and often between about 15 and 20 residues.

"Pharmaceutically acceptable" refers to a generally non-toxic, inert, and/or physiologically compatible composition.

15 A "pharmaceutical excipient" comprises a material such as an adjuvant, a carrier, pH-adjusting and buffering agents, tonicity adjusting agents, wetting agents, preservative, and the like.

A "primary anchor residue" is an amino acid at a specific position along a peptide sequence which is understood to provide a contact point between the immunogenic 20 peptide and the HLA molecule. One to three, usually two, primary anchor residues within a peptide of defined length generally defines a "motif" for an immunogenic peptide. These residues are understood to fit in close contact with peptide binding grooves of an HLA molecule, with their side chains buried in specific pockets of the binding grooves themselves. In one embodiment, the primary anchor residues are located 25 at position 2 (from the amino terminal position) and at the carboxyl terminal position of a 9-residue peptide epitope in accordance with the invention. The primary anchor positions for each motif and supermotif are set forth in Table 1. For example, analog peptides can be created by altering the presence or absence of particular residues in these primary anchor positions. Such analogs are used to modulate the binding affinity of a peptide 30 comprising a particular motif or supermotif.

"Promiscuous recognition" is where a distinct peptide is recognized by the same T cell clone in the context of various HLA molecules. Promiscuous recognition or binding is synonymous with cross-reactive binding.

A "protective immune response" or "therapeutic immune response" refers to a CTL and/or an HTL response to an antigen derived from an infectious agent or a tumor antigen, which prevents or at least partially arrests disease symptoms or progression. The immune response may also include an antibody response which has been facilitated by 5 the stimulation of helper T cells.

The term "residue" refers to an amino acid or amino acid mimetic incorporated into an oligopeptide by an amide bond or amide bond mimetic.

A "secondary anchor residue" is an amino acid at a position other than a primary anchor position in a peptide which may influence peptide binding. A secondary anchor 10 residue occurs at a significantly higher frequency amongst bound peptides than would be expected by random distribution of amino acids at one position. The secondary anchor residues are said to occur at "secondary anchor positions." A secondary anchor residue can be identified as a residue which is present at a higher frequency among high or intermediate affinity binding peptides, or a residue otherwise associated with high or 15 intermediate affinity binding. For example, analog peptides can be created by altering the presence or absence of particular residues in these secondary anchor positions. Such analogs are used to finely modulate the binding affinity of a peptide comprising a particular motif or supermotif.

A "subdominant epitope" is an epitope which evokes little or no response upon 20 immunization with whole antigens which comprise the epitope, but for which a response can be obtained by immunization with an isolated peptide, and this response (unlike the case of cryptic epitopes) is detected when whole protein is used to recall the response *in vitro* or *in vivo*.

A "supermotif" is a peptide binding specificity shared by HLA molecules encoded 25 by two or more HLA alleles. Preferably, a supermotif-bearing peptide is recognized with high or intermediate affinity (as defined herein) by two or more HLA antigens.

"Synthetic peptide" refers to a peptide that is man-made using such methods as chemical synthesis or recombinant DNA technology.

As used herein, a "vaccine" is a composition that contains one or more peptides of 30 the invention. There are numerous embodiments of vaccines in accordance with the invention, such as by a cocktail of one or more peptides; one or more epitopes of the invention comprised by a polyepitopic peptide; or nucleic acids that encode such peptides or polypeptides, e.g., a minigene that encodes a polyepitopic peptide. The "one or more peptides" can include, e.g., at least 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18,

19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 45, 50,  
55, 60, 65, 70, 75, 80, 85, 90, 95 or 100 or more peptides of the invention. The peptides  
or polypeptides can optionally be modified, such as by lipidation, addition of targeting or  
other sequences. HLA class I-binding peptides of the invention can be admixed with, or  
5 linked to, HLA class II-binding peptides, to facilitate activation of both cytotoxic T  
lymphocytes and helper T lymphocytes. Vaccines can also comprise peptide-pulsed  
antigen presenting cells, e.g., dendritic cells.

The nomenclature used to describe peptide compounds follows the conventional  
practice wherein the amino group is presented to the left (the N-terminus) and the  
10 carboxyl group to the right (the C-terminus) of each amino acid residue. When amino  
acid residue positions are referred to in a peptide epitope they are numbered in an amino  
to carboxyl direction with position one being the position closest to the amino terminal  
end of the epitope, or the peptide or protein of which it may be a part. In the formulae  
representing selected specific embodiments of the present invention, the amino- and  
15 carboxyl-terminal groups, although not specifically shown, are in the form they would  
assume at physiologic pH values, unless otherwise specified. In the amino acid structure  
formulae, each residue is generally represented by standard three letter or single letter  
designations. The L-form of an amino acid residue is represented by a capital single letter  
or a capital first letter of a three-letter symbol, and the D-form for those amino acids  
20 having D-forms is represented by a lower case single letter or a lower case three letter  
symbol. Glycine has no asymmetric carbon atom and is simply referred to as "Gly" or G.  
Symbols for the amino acids are shown below.

Single Letter Symbol	Three Letter Symbol	Amino Acids
A	Ala	Alanine
C	Cys	Cysteine
D	Asp	Aspartic Acid
E	Glu	Glutamic Acid
F	Phe	Phenylalanine
G	Gly	Glycine
H	His	Histidine
I	Ile	Isoleucine
K	Lys	Lysine
L	Leu	Leucine
M	Met	Methionine
N	Asn	Asparagine
P	Pro	Proline
Q	Gln	Glutamine
R	Arg	Arginine
S	Ser	Serine
T	Thr	Threonine
V	Val	Valine
W	Trp	Tryptophan
Y	Tyr	Tyrosine

#### IV.B. Stimulation of CTL and HTL responses

The mechanism by which T cells recognize antigens has been delineated during the past ten years. Based on our understanding of the immune system we have developed efficacious peptide epitope vaccine compositions that can induce a therapeutic or prophylactic immune response to HCV in a broad population. For an understanding of the value and efficacy of the claimed compositions, a brief review of immunology-related technology is provided.

A complex of an HLA molecule and a peptidic antigen acts as the ligand recognized by HLA-restricted T cells (Buus, S. *et al.*, *Cell* 47:1071, 1986; Babbitt, B. P. *et al.*, *Nature* 317:359, 1985; Townsend, A. and Bodmer, H., *Annu. Rev. Immunol.* 7:601,

1989; Germain, R. N., *Annu. Rev. Immunol.* 11:403, 1993). Through the study of single amino acid substituted antigen analogs and the sequencing of endogenously bound, naturally processed peptides, critical residues that correspond to motifs required for specific binding to HLA antigen molecules have been identified and are described herein and are set forth in Tables I, II, and III (see also, e.g., Southwood, *et al.*, *J. Immunol.* 160:3363, 1998; Rammensee, *et al.*, *Immunogenetics* 41:178, 1995; Rammensee *et al.*, SYFPEITHI, access via web at : <http://134.2.96.221/scripts.hlaserver.dll/home.htm>; Sette, A. and Sidney, J. *Curr. Opin. Immunol.* 10:478, 1998; Engelhard, V. H., *Curr. Opin. Immunol.* 6:13, 1994; Sette, A. and Grey, H. M., *Curr. Opin. Immunol.* 4:79, 1992; 10 Sinigaglia, F. and Hammer, J. *Curr. Biol.* 6:52, 1994; Ruppert *et al.*, *Cell* 74:929-937, 1993; Kondo *et al.*, *J. Immunol.* 155:4307-4312, 1995; Sidney *et al.*, *J. Immunol.* 157:3480-3490, 1996; Sidney *et al.*, *Human Immunol.* 45:79-93, 1996; Sette, A. and Sidney, J. *Immunogenetics*, in press, 1999).

Furthermore, x-ray crystallographic analysis of HLA-peptide complexes has 15 revealed pockets within the peptide binding cleft of HLA molecules which accommodate, in an allele-specific mode, residues borne by peptide ligands; these residues in turn determine the HLA binding capacity of the peptides in which they are present. (See, e.g., Madden, D.R. *Annu. Rev. Immunol.* 13:587, 1995; Smith, *et al.*, *Immunity* 4:203, 1996; Fremont *et al.*, *Immunity* 8:305, 1998; Stern *et al.*, *Structure* 2:245, 1994; Jones, E.Y. 20 *Curr. Opin. Immunol.* 9:75, 1997; Brown, J. H. *et al.*, *Nature* 364:33, 1993; Guo, H. C. *et al.*, *Proc. Natl. Acad. Sci. USA* 90:8053, 1993; Guo, H. C. *et al.*, *Nature* 360:364, 1992; Silver, M. L. *et al.*, *Nature* 360:367, 1992; Matsumura, M. *et al.*, *Science* 257:927, 1992; Madden *et al.*, *Cell* 70:1035, 1992; Fremont, D. H. *et al.*, *Science* 257:919, 1992; Saper, M. A. , Bjorkman, P. J. and Wiley, D. C., *J. Mol. Biol.* 219:277, 1991.)

25 Accordingly, the definition of class I and class II allele-specific HLA binding motifs, or class I or class II supermotifs allows identification of regions within a protein that have the potential of binding particular HLA antigen(s).

The present inventors have found that the correlation of binding affinity with 30 immunogenicity, which is disclosed herein, is an important factor to be considered when evaluating candidate peptides. Thus, by a combination of motif searches and HLA-peptide binding assays, candidates for epitope-based vaccines have been identified. After determining their binding affinity, additional confirmatory work can be performed to select, amongst these vaccine candidates, epitopes with preferred characteristics in terms of population coverage, antigenicity, and immunogenicity.

Various strategies can be utilized to evaluate immunogenicity, including:

- 1) Evaluation of primary T cell cultures from normal individuals (*see, e.g.*, Wentworth, P. A. *et al.*, *Mol. Immunol.* 32:603, 1995; Celis, E. *et al.*, *Proc. Natl. Acad. Sci. USA* 91:2105, 1994; Tsai, V. *et al.*, *J. Immunol.* 158:1796, 1997; Kawashima, I. *et al.*, *Human Immunol.* 59:1, 1998); This procedure involves the stimulation of peripheral blood lymphocytes (PBL) from normal subjects with a test peptide in the presence of antigen presenting cells *in vitro* over a period of several weeks. T cells specific for the peptide become activated during this time and are detected using, *e.g.*, a  $^{51}\text{Cr}$ -release assay involving peptide sensitized target cells.
- 10 2) Immunization of HLA transgenic mice (*see, e.g.*, Wentworth, P. A. *et al.*, *J. Immunol.* 26:97, 1996; Wentworth, P. A. *et al.*, *Int. Immunol.* 8:651, 1996; Alexander, J. *et al.*, *J. Immunol.* 159:4753, 1997); In this method, peptides in incomplete Freund's adjuvant are administered subcutaneously to HLA transgenic mice. Several weeks following immunization, splenocytes are removed and cultured *in vitro* in the presence of test peptide for approximately one week. Peptide-specific T cells are detected using, *e.g.*, a  $^{51}\text{Cr}$ -release assay involving peptide sensitized target cells and target cells expressing endogenously generated antigen.
- 15 3) Demonstration of recall T cell responses from immune individuals who have effectively been vaccinated, recovered from infection, and/or from chronically infected patients (*see, e.g.*, Rehermann, B. *et al.*, *J. Exp. Med.* 181:1047, 1995; Doolan, D. L. *et al.*, *Immunity* 7:97, 1997; Bertoni, R. *et al.*, *J. Clin. Invest.* 100:503, 1997; Threlkeld, S. C. *et al.*, *J. Immunol.* 159:1648, 1997; Diepolder, H. M. *et al.*, *J. Virol.* 71:6011, 1997). In applying this strategy, recall responses are detected by culturing PBL from subjects that have been naturally exposed to the antigen, for instance through infection, and thus 20 have generated an immune response "naturally", or from patients who were vaccinated against the infection. PBL from subjects are cultured *in vitro* for 1-2 weeks in the presence of test peptide plus antigen presenting cells (APC) to allow activation of "memory" T cells, as compared to "naive" T cells. At the end of the culture period, T cell 25 activity is detected using assays for T cell activity including  $^{51}\text{Cr}$  release involving peptide-sensitized targets, T cell proliferation, or lymphokine release.
- 30

The following describes the peptide epitopes and corresponding nucleic acids of the invention.

#### IV.C. Binding Affinity of Peptide Epitopes for HLA Molecules

The large degree of HLA polymorphism is an important factor to consider with the epitope-based approach to vaccine development. To address this factor, epitope selection including identification of peptides capable of binding at high or intermediate affinity to multiple HLA molecules is often utilized, most preferably these epitopes bind at high or intermediate affinity to two or more allele specific HLA molecules.

CTL-inducing peptides of interest for vaccine compositions preferably include those that have an  $IC_{50}$  or binding affinity value for class I HLA molecules of 500 nM or better (*i.e.*, the value is  $\leq$  500 nM). HTL-inducing peptides preferably include those that 10 have an  $IC_{50}$  or binding affinity value for class II HLA molecules of 1000 nM or better, (*i.e.*, the value is  $\leq$  1,000 nM). For example, peptide binding is assessed by testing the capacity of a candidate peptide to bind to a purified HLA molecule *in vitro*. Peptides exhibiting high or intermediate affinity are then considered for further analysis. Selected peptides are tested on other members of the supertype family. In preferred embodiments, 15 peptides that exhibit cross-reactive binding are then used in vaccines or in cellular screening analyses.

Higher HLA binding affinity is typically correlated with greater immunogenicity. Greater immunogenicity can be manifested in several different ways. Immunogenicity corresponds to whether an immune response is elicited at all, and to the vigor of any 20 particular response, as well as to the extent of a population in which a response is elicited. For example, a peptide might elicit an immune response in a diverse array of the population, yet in no instance produce a vigorous response. In accordance with these principles, close to 90% of high binding peptides have been found to be immunogenic, as contrasted with about 50% of the peptides which bind with intermediate affinity. 25 Moreover, higher binding affinity peptides leads to more vigorous immunogenic responses. As a result, less peptide is required to elicit a similar biological effect if a high affinity binding peptide is used. Thus, in preferred embodiments of the invention, high affinity binding epitopes are particularly useful.

The relationship between binding affinity for HLA class I molecules and 30 immunogenicity of discrete peptide epitopes on bound antigens has been determined for the first time in the art by the present inventors. The correlation between binding affinity and immunogenicity was analyzed in two different experimental approaches (*see, e.g.*, Sette, *et al.*, *J. Immunol.* 153:5586-5592, 1994). In the first approach, the

immunogenicity of potential epitopes ranging in HLA binding affinity over a 10,000-fold range was analyzed in HLA-A\*0201 transgenic mice. In the second approach, the antigenicity of approximately 100 different hepatitis B virus (HBV)-derived potential epitopes, all carrying A\*0201 binding motifs, was assessed by using PBL from acute hepatitis patients. Pursuant to these approaches, it was determined that an affinity threshold value of approximately 500 nM (preferably 50 nM or less) determines the capacity of a peptide epitope to elicit a CTL response. These data are true for class I binding affinity measurements for naturally processed peptides and for synthesized T cell epitopes. These data also indicate the important role of determinant selection in the shaping of T cell responses (see, e.g., Schaeffer *et al. Proc. Natl. Acad. Sci. USA* 86:4649-4653, 1989).

An affinity threshold associated with immunogenicity in the context of HLA class II DR molecules has also been delineated (see, e.g., Southwood *et al. J. Immunology* 160:3363-3373, 1998). In order to define a biologically significant threshold of DR binding affinity, a database of the binding affinities of 32 DR-restricted epitopes for their restricting element (*i.e.*, the HLA molecule that binds the motif) was compiled. In approximately half of the cases (15 of 32 epitopes), DR restriction was associated with high binding affinities, *i.e.* binding affinity values of 100 nM or less. In the other half of the cases (16 of 32), DR restriction was associated with intermediate affinity (binding affinity values in the 100-1000 nM range). In only one of 32 cases was DR restriction associated with an IC<sub>50</sub> of 1000 nM or greater. Thus, 1000 nM can be defined as an affinity threshold associated with immunogenicity in the context of DR molecules.

The binding affinity of peptides for HLA molecules can be determined as described in Example 1, below.

25

#### IV.D. Peptide Epitope Binding Motifs and Supermotifs

In the past few years evidence has accumulated to demonstrate that a large fraction of HLA class I and class II molecules can be classified into a relatively few supertypes, each characterized by largely overlapping peptide binding repertoires, and consensus structures of the main peptide binding pockets.

For HLA molecule pocket analyses, the residues comprising the B and F pockets of HLA class I molecules as described in crystallographic studies were analyzed (see, e.g., Guo, H. C. *et al., Nature* 360:364, 1992; Saper, M. A. , Bjorkman, P. J. and Wiley, D. C., *J. Mol. Biol.* 219:277, 1991; Madden, D. R., Garboczi, D. N. and Wiley, D. C.,

Cell 75:693, 1993; Parham, P., Adams, E. J., and Arnett, K. L., *Immunol. Rev.* 143:141, 1995). In these analyses, residues 9, 45, 63, 66, 67, 70, and 99 were considered to make up the B pocket; and the B pocket was deemed to determine the specificity for the amino acid residue in the second position of peptide ligands. Similarly, residues 77, 80, 81, and 5 116 were considered to determine the specificity of the F pocket; the F pocket was deemed to determine the specificity for the C-terminal residue of a peptide ligand bound by the HLA class I molecule.

Through the study of single amino acid substituted antigen analogs and the sequencing of endogenously bound, naturally processed peptides, critical residues 10 required for allele-specific binding to HLA molecules have been identified. The presence of these residues correlates with binding affinity for HLA molecules. The identification of motifs and/or supermotifs that correlate with high and intermediate affinity binding is an important issue with respect to the identification of immunogenic peptide epitopes for the inclusion in a vaccine. Kast *et al.* (*J. Immunol.* 152:3904-3912, 1994) have shown 15 that motif-bearing peptides account for 90% of the epitopes that bind to allele-specific HLA class I molecules. In this study all possible peptides of 9 amino acids in length and overlapping by eight amino acids (240 peptides), which cover the entire sequence of the E6 and E7 proteins of human papillomavirus type 16, were evaluated for binding to five allele-specific HLA molecules that are expressed at high frequency among different 20 ethnic groups. This unbiased set of peptides allowed an evaluation of the predictive value of HLA class I motifs. From the set of 240 peptides, 22 peptides were identified that bound to an allele-specific HLA molecule with high or intermediate affinity. Of these 22 peptides, 20 (*i.e.* 91%) were motif-bearing. Thus, this study demonstrates the value of motifs for the identification of peptide epitopes for inclusion in a vaccine: application of 25 motif-based identification techniques eliminates screening of 90% of the potential epitopes in a target antigen protein sequence.

Such peptide epitopes are identified in the Tables described below.

Peptides of the present invention may also comprise epitopes that bind to MHC 30 class II DR molecules. A greater degree of heterogeneity in both size and binding frame position of the motif, relative to the N and C termini of the peptide, exists for class II peptide ligands. This increased heterogeneity of HLA class II peptide ligands is due to the structure of the binding groove of the HLA class II molecule which, unlike its class I counterpart, is open at both ends. Crystallographic analysis of HLA class II DRB\*0101-peptide complexes showed that the major energy of binding is contributed by peptide

residues complexed with complementary pockets on the DRB\*0101 molecules. An important anchor residue engages the deepest hydrophobic pocket (*see, e.g.*, Madden, D.R. *Ann. Rev. Immunol.* 13:587, 1995) and is referred to as position 1 (P1). P1 may represent the N-terminal residue of a class II binding peptide epitope, but more typically 5 is flanked towards the N-terminus by one or more residues. Other studies have also pointed to an important role for the peptide residue in the 6<sup>th</sup> position towards the C-terminus, relative to P1, for binding to various DR molecules.

Thus, peptides of the present invention are identified by any one of several HLA-specific amino acid motifs (*see, e.g.*, Tables I-III). If the presence of the motif 10 corresponds to the ability to bind several allele-specific HLA antigens, it is referred to as a supermotif. The HLA molecules that bind to peptides that possess a particular amino acid supermotif are collectively referred to as an HLA "supertype."

The peptide motifs and supermotifs described below, and summarized in Tables I-III, provide guidance for the identification and use of peptide epitopes in accordance with 15 the invention.

Examples of peptide epitopes bearing a respective supermotif or motif are included in Tables as designated in the description of each motif or supermotif below. The Tables include a binding affinity ratio listing for some of the peptide epitopes. The ratio may be converted to IC<sub>50</sub> by using the following formula: IC<sub>50</sub> of the standard 20 peptide/ratio = IC<sub>50</sub> of the test peptide (*i.e.*, the peptide epitope). The IC<sub>50</sub> values of standard peptides used to determine binding affinities for Class I peptides are shown in Table IV. The IC<sub>50</sub> values of standard peptides used to determine binding affinities for Class II peptides are shown in Table V. The peptides used as standards for the binding assays described herein are examples of standards; alternative standard peptides can also 25 be used when performing such an analysis.

To obtain the peptide epitope sequences listed in each Table, protein sequence data from fourteen HCV isolates were evaluated for the presence of the designated supermotif or motif. The fourteen strains include HPCCGAA, HPCPLYPRE, HCV-H-CMR, HCV-J1, HPCGENANTI, HPCGENOM, HPCHUMR, HPCJCG, HPCJTA, HCV-30 J483, HCV-JK1, HCV-N, HPCPOLP, and HCV-J8. Peptide epitopes were additionally evaluated on the basis of their conservancy among these fourteen strains. A criterion for conservancy requires that the entire sequence of an HLA class I binding peptide be totally conserved in 79% of the sequences available for a specific protein. Similarly, a criterion for conservancy requires that the entire 9-mer core region of an HLA class II binding

peptide be totally conserved in 79% of the sequences available for a specific protein. The percent conservancy of the selected peptide epitopes is indicated on the Tables. The frequency, *i.e.* the number of strains of the fourteen strains in which the totally conserved peptide sequence was identified, is also shown. The "position" column in the Tables 5 designates the amino acid position of the HCV polyprotein that corresponds to the first amino acid residue of the epitope. The "number of amino acids" indicates the number of residues in the epitope sequence.

#### **HLA Class I Motifs Indicative of CTL Inducing Peptide Epitopes:**

10 The primary anchor residues of the HLA class I peptide epitope supermotifs and motifs delineated below are summarized in Table I. The HLA class I motifs set out in Table I(a) are those most particularly relevant to the invention claimed here. Primary and secondary anchor positions are summarized in Table II. Allele-specific HLA molecules that comprise HLA class I supertype families are listed in Table VI.

15

##### **IV.D.1. HLA-A1 supermotif**

The HLA-A1 supermotif is characterized by the presence in peptide ligands of a small (T or S) or hydrophobic (L, I, V, or M) primary anchor residue in position 2, and an aromatic (Y, F, or W) primary anchor residue at the C-terminal position of the epitope.

20 The corresponding family of HLA molecules that bind to the A1 supermotif (*i.e.*, the HLA-A1 supertype) includes at least A\*0101, A\*2601, A\*2602, A\*2501, and A\*3201 (*see, e.g.*, DiBrino, M. *et al.*, *J. Immunol.* 151:5930, 1993; DiBrino, M. *et al.*, *J. Immunol.* 152:620, 1994; Kondo, A. *et al.*, *Immunogenetics* 45:249, 1997). Other allele-specific HLA molecules predicted to be members of the A1 superfamily are shown in 25 Table VI. Peptides binding to each of the individual HLA proteins can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the A1 supermotif are set forth in Table VII.

30 **IV.D.2. HLA-A2 supermotif**

Primary anchor specificities for allele-specific HLA-A2.1 molecules (Falk *et al.*, *Nature* 351:290-296, 1991; Hunt *et al.*, *Science* 255:1261-1263, 1992; Parker *et al.*, *J. Immunol.* 149:3580-3587, 1992) and cross-reactive binding within the HLA A2 family (Fruci *et al.*, *Human Immunol.* 38:187-192, 1993; Tanigaki *et al.*, *Human Immunol.*

39:155-162, 1994) have been described. The present inventors have defined additional primary anchor residues that determine cross-reactive binding to multiple allele-specific HLA A2 molecules (Ruppert *et al.*, *Cell* 74:929-937, 1993; Del Guercio *et al.*, *J. Immunol.* 154:685-693, 1995; Kast *et al.*, *J. Immunol.* 152:3904-3912, 1994). The HLA-  
5 A2 supermotif comprises peptide ligands with L, I, V, M, A, T, or Q as a primary anchor residue at position 2 and L, I, V, M, A, or T as a primary anchor residue at the C-terminal position of the epitope.

The corresponding family of HLA molecules (*i.e.*, the HLA-A2 supertype that binds these peptides) is comprised of at least: A\*0201, A\*0202, A\*0203, A\*0204,  
10 A\*0205, A\*0206, A\*0207, A\*0209, A\*0214, A\*6802, and A\*6901. Other allele-specific HLA molecules predicted to be members of the A2 superfamily are shown in Table VI. As explained in detail below, binding to each of the individual allele-specific HLA molecules can be modulated by substitutions at the primary anchor and/or secondary anchor positions, preferably choosing respective residues specified for the  
15 supermotif.

Peptide epitopes that comprise an A2 supermotif are set forth in Table VIII. The motifs comprising the primary anchor residues V, A, T, or Q at position 2 and L, I, V, A, or T at the C-terminal position are those most particularly relevant to the invention claimed herein.

20

#### IV.D.3. HLA-A3 supermotif

The HLA-A3 supermotif is characterized by the presence in peptide ligands of A, L, I, V, M, S, or, T as a primary anchor at position 2, and a positively charged residue, R or K, at the C-terminal position of the epitope (*e.g.*, in position 9 of 9-mers). Exemplary members of the corresponding family of HLA molecules (the HLA-A3 supertype) that bind the A3 supermotif include at least A\*0301, A\*1101, A\*3101, A\*3301, and A\*6801. Other allele-specific HLA molecules predicted to be members of the A3 superfamily are shown in Table VI. As explained in detail below, peptide binding to each of the individual allele-specific HLA proteins can be modulated by substitutions of amino acids at the primary and/or secondary anchor positions of the peptide, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the A3 supermotif are set forth in Table IX.

**IV.D.4. HLA-A24 supermotif**

The HLA-A24 supermotif is characterized by the presence in peptide ligands of an aromatic (F, W, or Y) or hydrophobic aliphatic (L, I, V, M, or T) residue as a primary anchor in position 2, and Y, F, W, L, I, or M as primary anchor at the C-terminal position 5 of the epitope. The corresponding family of HLA molecules that bind to the A24 supermotif (*i.e.*, the A24 supertype) includes at least A\*2402, A\*3001, and A\*2301. Other allele-specific HLA molecules predicted to be members of the A24 superfamily are shown in Table VI. Peptide binding to each of the allele-specific HLA molecules can be modulated by substitutions at primary and/or secondary anchor positions, preferably 10 choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the A24 supermotif are set forth in Table X.

**IV.D.5. HLA-B7 supermotif**

The HLA-B7 supermotif is characterized by peptides bearing proline in position 2 15 as a primary anchor, and a hydrophobic or aliphatic amino acid (L, I, V, M, A, F, W, or Y) as the primary anchor at the C-terminal position of the epitope. The corresponding family of HLA molecules that bind the B7 supermotif (*i.e.*, the HLA-B7 supertype) is comprised of at least twenty six HLA-B proteins including: B\*0702, B\*0703, B\*0704, B\*0705, B\*1508, B\*3501, B\*3502, B\*3503, B\*3504, B\*3505, B\*3506, B\*3507, 20 B\*3508, B\*5101, B\*5102, B\*5103, B\*5104, B\*5105, B\*5301, B\*5401, B\*5501, B\*5502, B\*5601, B\*5602, B\*6701, and B\*7801 (*see, e.g.*, Sidney, *et al.*, *J. Immunol.* 154:247, 1995; Barber, *et al.*, *Curr. Biol.* 5:179, 1995; Hill, *et al.*, *Nature* 360:434, 1992; Rammensee, *et al.*, *Immunogenetics* 41:178, 1995). Other allele-specific HLA molecules predicted to be members of the B7 superfamily are shown in Table VI. As explained in 25 detail below, peptide binding to each of the individual allele-specific HLA proteins can be modulated by substitutions at the primary and/or secondary anchor positions of the peptide, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the B7 supermotif are set forth in Table XI.

**IV.D.6. HLA-B27 supermotif**

The HLA-B27 supermotif is characterized by the presence in peptide ligands of a positively charged (R, H, or K) residue as a primary anchor at position 2, and a hydrophobic (F, Y, L, W, M, I, A, or V) residue as a primary anchor at the C-terminal position of the epitope. Exemplary members of the corresponding family of HLA

molecules that bind to the B27 supermotif (*i.e.*, the B27 supertype) include at least B\*1401, B\*1402, B\*1509, B\*2702, B\*2703, B\*2704, B\*2705, B\*2706, B\*3801, B\*3901, B\*3902, and B\*7301. Other allele-specific HLA molecules predicted to be members of the B27 superfamily are shown in Table VI. Peptide binding to each of the 5 allele-specific HLA molecules can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the B27 supermotif are set forth in Table XII.

10 **IV.D.7. HLA-B44 supermotif**

The HLA-B44 supermotif is characterized by the presence in peptide ligands of negatively charged (D or E) residues as a primary anchor in position 2, and hydrophobic residues (F, W, Y, L, I, M, V, or A) as a primary anchor at the C-terminal position of the epitope. Exemplary members of the corresponding family of HLA molecules that bind to 15 the B44 supermotif (*i.e.*, the B44 supertype) include at least: B\*1801, B\*1802, B\*3701, B\*4001, B\*4002, B\*4006, B\*4402, B\*4403, and B\*4006. Peptide binding to each of the allele-specific HLA molecules can be modulated by substitutions at primary and/or secondary anchor positions; preferably choosing respective residues specified for the supermotif.

20

**IV.D.8. HLA-B58 supermotif**

The HLA-B58 supermotif is characterized by the presence in peptide ligands of a small aliphatic residue (A, S, or T) as a primary anchor residue at position 2, and an aromatic or hydrophobic residue (F, W, Y, L, I, V, M, or A) as a primary anchor residue 25 at the C-terminal position of the epitope. Exemplary members of the corresponding family of HLA molecules that bind to the B58 supermotif (*i.e.*, the B58 supertype) include at least: B\*1516, B\*1517, B\*5701, B\*5702, and B\*5801. Other allele-specific HLA molecules predicted to be members of the B58 superfamily are shown in Table VI. Peptide binding to each of the allele-specific HLA molecules can be modulated by 30 substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the B58 supermotif are set forth in Table XIII.

**IV.D.9. HLA-B62 supermotif**

The HLA-B62 supermotif is characterized by the presence in peptide ligands of the polar aliphatic residue Q or a hydrophobic aliphatic residue (L, V, M, I, or P) as a primary anchor in position 2, and a hydrophobic residue (F, W, Y, M, I, V, L, or A) as a primary anchor at the C-terminal position of the epitope. Exemplary members of the corresponding family of HLA molecules that bind to the B62 supermotif (*i.e.*, the B62 supertype) include at least: B\*1501, B\*1502, B\*1513, and B5201. Other allele-specific HLA molecules predicted to be members of the B62 superfamily are shown in Table VI. Peptide binding to each of the allele-specific HLA molecules can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the supermotif.

Peptide epitopes that comprise the B62 supermotif are set forth in Table XIV.

**IV.D.10. HLA-A1 motif**

The HLA-A1 motif is characterized by the presence in peptide ligands of T, S, or M as a primary anchor residue at position 2 and the presence of Y as a primary anchor residue at the C-terminal position of the epitope. An alternative allele-specific A1 motif is characterized by a primary anchor residue at position 3 rather than position 2. This motif is characterized by the presence of D, E, A, or S as a primary anchor residue in position 3, and a Y as a primary anchor residue at the C-terminal position of the epitope. Peptide binding to HLA A1 can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the motif.

Peptide epitopes that comprise either A1 motif are set forth in Table XV. The epitopes comprising T, S, or M at position 2 and Y at the C-terminal position are also included in the listing of HLA-A1 supermotif-bearing peptide epitopes listed in Table VII.

**IV.D.11. HLA-A\*0201 motif**

An HLA-A2\*0201 motif was first determined to be characterized by the presence in peptide ligands of L or M as a primary anchor residue in position 2, and L or V as a primary anchor residue at the C-terminal position of a 9-residue peptide (Falk *et al.*, *Nature* 351:290-296, 1991). The A\*0201 motif was also determined to further comprise an I at position 2 and I or A at the C-terminal position of a nine amino acid peptide (Hunt

*et al.*, *Science* 255:1261-1263, March 6, 1992; Parker *et al.*, *J. Immunol.* 149:3580-3587, 1992). Subsequently, the A\*0201 allele-specific motif has been defined by the present inventors to additionally comprise V, A, T, or Q as a primary anchor residue at position 2, and M as a primary anchor residue at the C-terminal position of the epitope.

5 Additionally, the A\*0201 allele-specific motif has been found to comprise a T at the C-terminal position (Kast *et al.*, *J. Immunol.* 152:3904-3912, 1994). Thus, the HLA-A\*0201 motif comprises peptide ligands with L, I, V, M, A, T, or Q as primary anchor residues at position 2 and L, I, V, M, A, or T as a primary anchor residue at the C-terminal position of the epitope. The preferred and tolerated residues that characterize the 10 primary anchor positions of the HLA-A\*0201 motif are identical to the residues describing the A2 supermotif. (For reviews of relevant data, *see, e.g.*, Del Guercio *et al.*, *J. Immunol.* 154:685-693, 1995; Ruppert *et al.*, *Cell* 74:929-937, 1993; Sidney *et al.*, *Immunol. Today* 17:261-266, 1996; Sette and Sidney, *Curr. Opin. in Immunol.* 10:478-482, 1998). Secondary anchor residues that characterize the A\*0201 motif have 15 additionally been defined as disclosed herein. These are disclosed in Table II. Peptide binding to HLA-A\*0201 molecules can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the motif.

Peptide epitopes that comprise an A\*0201 motif are set forth in Table VIII. The 20 A\*0201 motifs comprising the primary anchor residues V, A, T, or Q at position 2 and L, I, V, A, or T at the C-terminal position are those most particularly relevant to the invention claimed herein.

#### IV.D.12. HLA-A3 motif

25 The HLA-A3 motif is characterized by the presence in peptide ligands of L, M, V, I, S, A, T, F, C, G, or D as a primary anchor residue at position 2, and the presence of K, Y, R, H, F, or A as a primary anchor residue at the C-terminal position of the epitope. Peptide binding to HLA-A3 can be modulated by substitutions at primary and/or 30 secondary anchor positions, preferably choosing respective residues specified for the motif.

The A3 supermotif primary anchor residues comprise a subset of the A3- and A11-allele specific motif primary anchor residues. Peptide epitopes that comprise the A3 motif are set forth in Table XVI. Those peptide epitopes that also comprise the A3 supermotif are also listed in Table IX.

**IV.D.13. HLA-A11 motif**

The HLA-A11 motif is characterized by the presence in peptide ligands of V, T, M, L, I, S, A, G, N, C, D, or F as a primary anchor residue in position 2, and K, R, Y, or 5 H as a primary anchor residue at the C-terminal position of the epitope. Peptide binding to HLA-A11 can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the motif.

Peptide epitopes that comprise the A11 motif are set forth in Table XVII; peptide epitopes comprising the A3 allele-specific motif are also present in this Table because of 10 the overlap between the A3 and A11 motif primary anchor specificities. Further, those peptide epitopes that comprise the A3 supermotif are also listed in Table IX.

**IV.D.14. HLA-A24 motif**

The HLA-A24 motif is characterized by the presence in peptide ligands of Y, F, 15 W, or M as a primary anchor residue in position 2, and F, L, I, or W as a primary anchor residue at the C-terminal position of the epitope. Peptide binding to HLA-A24 molecules can be modulated by substitutions at primary and/or secondary anchor positions; preferably choosing respective residues specified for the motif.

Peptide epitopes that comprise the A24 motif are set forth in Table XVIII. These 20 epitopes are also listed in Table X, which sets forth HLA-A24-supermotif-bearing peptide epitopes, as the primary anchor residues characterizing the A24 allele-specific motif comprise a subset of the A24 supermotif primary anchor residues.

**HLA Class II Binding Motifs**

25 The primary and secondary anchor residues of the HLA class II peptide epitope supermotifs and motifs delineated below are summarized in Table III.

**IV.D.15. HLA DR-1-4-7 supermotif**

Motifs have also been identified for peptides that bind to three common HLA 30 class II allele-specific HLA molecules: HLA DRB1\*0401, DRB1\*0101, and DRB1\*0701. Collectively, the common residues from these motifs delineate the HLA DR-1-4-7 supermotif. Peptides that bind to these DR molecules carry a supermotif characterized by a large aromatic or hydrophobic residue (Y, F, W, L, I, V, or M) as a primary anchor residue in position 1, and a small, non-charged residue (S, T, C, A, P, V,

I, L, or M) as a primary anchor residue in position 6 of a 9-mer core region. Allele-specific secondary effects and secondary anchors for each of these HLA types have also been identified. These are set forth in Table III. Peptide binding to HLA- DRB1\*0401, DRB1\*0101, and/or DRB1\*0701 can be modulated by substitutions at primary and/or 5 secondary anchor positions, preferably choosing respective residues specified for the supermotif.

Conserved peptide epitopes *i.e.*, conserved in  $\geq 79\%$  ( $\geq 11/14$ ) of the HCV strains used for the present analysis, may be described as corresponding to epitopes containing a nine residue core comprising the DR-1-4-7 supermotif, and in which the 9 residue core is 10 conserved in  $\geq 79\%$  (wherein position 1 of the motif is at position 1 of the nine residue core). Conserved 9-mer core regions are set forth in Table XIXa. Respective exemplary peptide epitopes of 15 amino acid residues in length, each of which comprise a conserved nine residue core, are also shown in section "a" of the Table. Cross-reactive binding data for exemplary 15-residue supermotif-bearing peptides are shown in Table XIXb.

15

#### IV.D.16. HLA DR3 motifs

Two alternative motifs (*i.e.*, submotifs) characterize peptide epitopes that bind to HLA-DR3 molecules. In the first motif (submotif DR3A) a large, hydrophobic residue (L, I, V, M, F, or Y) is present in anchor position 1 of a 9-mer core, and D is present as an 20 anchor at position 4, towards the carboxyl terminus of the epitope. As in other class II motifs, core position 1 may or may not occupy the peptide N-terminal position.

The alternative DR3 submotif provides for lack of the large, hydrophobic residue at anchor position 1, and/or lack of the negatively charged or amide-like anchor residue at position 4, by the presence of a positive charge at position 6 towards the carboxyl 25 terminus of the epitope. Thus, for the alternative allele-specific DR3 motif (submotif DR3B): L, I, V, M, F, Y, A, or Y is present at anchor position 1; D, N, Q, E, S, or T is present at anchor position 4; and K, R, or H is present at anchor position 6. Peptide binding to HLA-DR3 can be modulated by substitutions at primary and/or secondary anchor positions, preferably choosing respective residues specified for the motif.

30

Conserved 9-mer core regions (*i.e.*, those sequences that are conserved in at least 79% of the 14 HCV strains used for the analysis) corresponding to a nine residue sequence comprising the DR3A submotif (wherein position 1 of the motif is at position 1 of the nine residue core) are set forth in Table XXa. Respective exemplary peptide

epitopes of 15 amino acid residues in length, each of which comprise a conserved nine residue core, are also shown in Table XXa. Table XXb shows binding data of exemplary DR3 submotif A-bearing peptides.

Conserved 9-mer core regions (*i.e.*, those that are at least 79% conserved in the 14 HCV strains used for the analysis) comprising the DR3B submotif and respective exemplary 15-mer peptides comprising the DR3 submotif-B epitope are set forth in Table XXc. Table XXd shows binding data of exemplary DR3 submotif B-bearing peptides.

Each of the HLA class I or class II peptide epitopes set out in the Tables herein are deemed singly to be an inventive aspect of this application. Further, it is also an inventive aspect of this application that each peptide epitope may be used in combination with any other peptide epitope.

#### IV.E. Enhancing Population Coverage of the Vaccine

Vaccines that have broad population coverage are preferred because they are more commercially viable and generally applicable to the most people. Broad population coverage can be obtained using the peptides of the invention (and nucleic acid compositions that encode such peptides) through selecting peptide epitopes that bind to HLA alleles which, when considered in total, are present in most of the population. Table XXI lists the overall frequencies of the HLA class I supertypes in various ethnicities (Table XXIa) and the combined population coverage achieved by the A2-, A3-, and B7-supertypes (Table XXIb). The A2-, A3-, and B7 supertypes are each present on the average of over 40% in each of these five major ethnic groups. Coverage in excess of 80% is achieved with a combination of these supermotifs. These results suggest that effective and non-ethnically biased population coverage is achieved upon use of a limited number of cross-reactive peptides. Although the population coverage reached with these three main peptide specificities is high, coverage can be expanded to reach 95% population coverage and above, and more easily achieve truly multispecific responses upon use of additional supermotif or allele-specific motif bearing peptides.

The B44-, A1-, and A24-supertypes are present, on average, in a range from 25% to 40% in these major ethnic populations (Table XXIa). While less prevalent overall, the B27-, B58-, and B62 supertypes are each present with a frequency >25% in at least one major ethnic group (Table XXIa). Table XXIb summarizes the estimated prevalence of combinations of HLA supertypes that have been identified in five major ethnic groups.

The incremental coverage obtained by the inclusion of A1,- A24-, and B44-supertypes to the A2, A3, and B7 coverage, or all of the supertypes described herein, is shown.

The data presented herein, together with the previous definition of the A2-, A3-, and B7-supertypes, indicates that all antigens, with the possible exception of A29, B8, and B46, can be classified into a total of nine HLA supertypes. By including epitopes from the six most frequent supertypes, an average population coverage of 99% is obtained for five major ethnic groups..

#### IV.F. Immune Response-Stimulating Peptide Analogs

10 In general, CTL and HTL responses are not directed against all possible epitopes. Rather, they are restricted to a few "immunodominant" determinants (Zinkernagel, *et al.*, *Adv. Immunol.* 27:5159, 1979; Bennink, *et al.*, *J. Exp. Med.* 168:19351939, 1988; Rawle, *et al.*, *J. Immunol.* 146:3977-3984, 1991). It has been recognized that immunodominance (Benacerraf, *et al.*, *Science* 175:273-279, 1972) could be explained by either the ability of 15 a given epitope to selectively bind a particular HLA protein (determinant selection theory) (Vitiello, *et al.*, *J. Immunol.* 131:1635, 1983); Rosenthal, *et al.*, *Nature* 267:156-158, 1977), or to be selectively recognized by the existing TCR (T cell receptor) specificities (repertoire theory) (Klein, J., *IMMUNOLOGY, THE SCIENCE OF SELFNONSELF DISCRIMINATION*, John Wiley & Sons, New York, pp. 270-310, 1982). It has been 20 demonstrated that additional factors, mostly linked to processing events, can also play a key role in dictating, beyond strict immunogenicity, which of the many potential determinants will be presented as immunodominant (Sercarz, *et al.*, *Annu. Rev. Immunol.* 11:729-766, 1993).

25 The concept of dominance and subdominance is relevant to immunotherapy of both infectious diseases and cancer. For example, in the course of chronic viral disease, recruitment of subdominant epitopes can be important for successful clearance of the infection, especially if dominant CTL or HTL specificities have been inactivated by functional tolerance, suppression, mutation of viruses and other mechanisms (Franco, *et al.*, *Curr. Opin. Immunol.* 7:524-531, 1995). In the case of cancer and tumor antigens, 30 CTLs recognizing at least some of the highest binding affinity peptides might be functionally inactivated. Lower binding affinity peptides are preferentially recognized at these times, and may therefore be preferred in therapeutic or prophylactic anti-cancer vaccines.

In particular, it has been noted that a significant number of epitopes derived from known non-viral tumor associated antigens (TAA) bind HLA class I with intermediate affinity ( $IC_{50}$  in the 50-500 nM range). For example, it has been found that 8 of 15 known TAA peptides recognized by tumor infiltrating lymphocytes (TIL) or CTL bound in the 50-500 nM range. (These data are in contrast with estimates that 90% of known viral antigens were bound by HLA class I molecules with  $IC_{50}$  of 50 nM or less, while only approximately 10% bound in the 50-500 nM range (Sette, *et al.*, *J. Immunol.*, 153:558-5592, 1994). In the cancer setting this phenomenon is probably due to elimination or functional inhibition of the CTL recognizing several of the highest binding peptides, presumably because of T cell tolerization events.

Without intending to be bound by theory, it is believed that because T cells to dominant epitopes may have been clonally deleted, selecting subdominant epitopes may allow existing T cells to be recruited, which will then lead to a therapeutic or prophylactic response. However, the binding of HLA molecules to subdominant epitopes is often less vigorous than to dominant ones. Accordingly, there is a need to be able to modulate the binding affinity of particular immunogenic epitopes for one or more HLA molecules, and thereby to modulate the immune response elicited by the peptide, for example to prepare analog peptides which elicit a more vigorous response. This ability would greatly enhance the usefulness of peptide-based vaccines and therapeutic agents.

Although peptides with suitable cross-reactivity among all alleles of a superfamily are identified by the screening procedures described above, cross-reactivity is not always as complete as possible, and in certain cases procedures to increase cross-reactivity of peptides can be useful; moreover, such procedures can also be used to modify other properties of the peptides such as binding affinity or peptide stability. Having established the general rules that govern cross-reactivity of peptides for HLA alleles within a given motif or supermotif, modification (*i.e.*, analoging) of the structure of peptides of particular interest in order to achieve broader (or otherwise modified) HLA binding capacity can be performed. More specifically, peptides which exhibit the broadest cross-reactivity patterns, can be produced in accordance with the teachings herein. The present concepts related to analog generation are set forth in greater detail in co-pending U.S.S.N. 09/226,775 filed 1/6/99.

In brief, the strategy employed utilizes the motifs or supermotifs which correlate with binding to certain HLA molecules. The motifs or supermotifs are defined by having primary anchors, and in many cases secondary anchors. Analog peptides can be created

by substituting amino acid residues at primary anchor, secondary anchor, or at primary and secondary anchor positions. Generally, analogs are made for peptides that already bear a motif or supermotif. Preferred secondary anchor residues of supermotifs and motifs that have been defined for HLA class I and class II binding peptides are shown in Tables II and III, respectively.

For a number of the motifs or supermotifs in accordance with the invention, residues are defined which are deleterious to binding to allele-specific HLA molecules or members of HLA supertypes that bind the respective motif or supermotif (Tables II and III). Accordingly, removal of such residues that are detrimental to binding can be performed in accordance with the present invention. For example, in the case of the A3 supertype, when all peptides that have such deleterious residues are removed from the population of analyzed peptides, the incidence of cross-reactivity increases from 22% to 37% (see, e.g., Sidney, J. et al., *Hu. Immunol.* 45:79, 1996). Thus, one strategy to improve the cross-reactivity of peptides within a given supermotif is simply to delete one or more of the deleterious residues present within a peptide and substitute a small “neutral” residue such as Ala (that may not influence T cell recognition of the peptide). An enhanced likelihood of cross-reactivity is expected if, together with elimination of detrimental residues within a peptide, “preferred” residues associated with high affinity binding to an allele-specific HLA molecule or to multiple HLA molecules within a superfamily are inserted.

To ensure that an analog peptide, when used as a vaccine, actually elicits a CTL response to the native epitope *in vivo* (or, in the case of class II epitopes, elicits helper T cells that cross-react with the wild type peptides), the analog peptide may be used to immunize T cells *in vitro* from individuals of the appropriate HLA allele. Thereafter, the immunized cells' capacity to induce lysis of wild type peptide sensitized target cells is evaluated. It will be desirable to use as antigen presenting cells, cells that have been either infected, or transfected with the appropriate genes, or, in the case of class II epitopes only, cells that have been pulsed with whole protein antigens, to establish whether endogenously produced antigen is also recognized by the relevant T cells.

Another embodiment of the invention is to create analogs of weak binding peptides, to thereby ensure adequate numbers of cross-reactive cellular binders. Class I binding peptides exhibiting binding affinities of 500-5000 nM, and carrying an acceptable but suboptimal primary anchor residue at one or both positions can be “fixed” by

substituting preferred anchor residues in accordance with the respective supertype. The analog peptides can then be tested for crossbinding activity.

Another embodiment for generating effective peptide analogs involves the substitution of residues that have an adverse impact on peptide stability or solubility in, 5 *e.g.*, a liquid environment. This substitution may occur at any position of the peptide epitope. For example, a cysteine (C) can be substituted out in favor of  $\alpha$ -amino butyric acid. Due to its chemical nature, cysteine has the propensity to form disulfide bridges and sufficiently alter the peptide structurally so as to reduce binding capacity. Substituting  $\alpha$ -amino butyric acid for C not only alleviates this problem, but actually improves binding 10 and crossbinding capability in certain instances (*see, e.g.*, the review by Sette *et al.*, In: Persistent Viral Infections, Eds. R. Ahmed and I. Chen, John Wiley & Sons, England, 1999). Substitution of cysteine with  $\alpha$ -amino butyric acid may occur at any residue of a peptide epitope, *i.e.* at either anchor or non-anchor positions.

15 Representative analog peptides are set forth in Table XXII. The Table indicates the length and sequence of the analog peptide as well as the motif or supermotif, if appropriate. The information in the "Fixed Nomenclature" column indicates the residues substituted at the indicated position numbers for the respective analog.

#### IV.G. Computer Screening of Protein Sequences from Disease-Related Antigens for 20 Supermotif- or Motif-Bearing Peptides

In order to identify supermotif- or motif-bearing epitopes in a target antigen, a native protein sequence, *e.g.*, a tumor-associated antigen, or sequences from an infectious organism, or a donor tissue for transplantation, is screened using a means for computing, such as an intellectual calculation or a computer, to determine the presence of a 25 supermotif or motif within the sequence. The information obtained from the analysis of native peptide can be used directly to evaluate the status of the native peptide or may be utilized subsequently to generate the peptide epitope.

Computer programs that allow the rapid screening of protein sequences for the occurrence of the subject supermotifs or motifs are encompassed by the present 30 invention; as are programs that permit the generation of analog peptides. These programs are implemented to analyze any identified amino acid sequence or operate on an unknown sequence and simultaneously determine the sequence and identify motif-bearing epitopes thereof; analogs can be simultaneously determined as well. Generally, the identified sequences will be from a pathogenic organism or a tumor-associated peptide. For

example, the target molecules considered herein include, without limitation, the core, S, E1, NS1/E2, NS2, NS3, NS4, and NS5 regions of HCV.

In cases where the sequence of multiple variants of the same target protein are available, peptides may also be selected on the basis of their conservancy. A presently preferred criterion for conservancy defines that the entire sequence of an HLA class I binding peptide or the entire 9-mer core of a class II binding peptide, be totally (*i.e.*, 100%) conserved in at least 79% of the sequences evaluated for a specific protein. This definition of conservancy has been employed herein; although, as appreciated by those in the art, lower or higher degrees of conservancy can be employed as appropriate for a given antigenic target.

It is important that the selection criteria utilized for prediction of peptide binding are as accurate as possible, to correlate most efficiently with actual binding. Prediction of peptides that bind, for example, to HLA-A\*0201, on the basis of the presence of the appropriate primary anchors, is positive at about a 30% rate (see, *e.g.*, Ruppert, J. *et al.* 15 *Cell* 74:929, 1993). However, by extensively analyzing peptide-HLA binding data disclosed herein, data in related patent applications, and data in the art, the present inventors have developed a number of allele-specific polynomial algorithms that dramatically increase the predictive value over identification on the basis of the presence of primary anchor residues alone. These algorithms take into account not only the 20 presence or absence of primary anchors, but also consider the positive or deleterious presence of secondary anchor residues (to account for the impact of different amino acids at different positions). The algorithms are essentially based on the premise that the overall affinity (or  $\Delta G$ ) of peptide-HLA interactions can be approximated as a linear polynomial function of the type:

$$\Delta G = a_{1i} \times a_{2i} \times a_{3i} \dots \times a_{ni}$$

where  $a_{ji}$  is a coefficient that represents the effect of the presence of a given amino acid ( $j$ ) at a given position ( $i$ ) along the sequence of a peptide of  $n$  amino acids. An important assumption of this method is that the effects at each position are essentially independent of each other. This assumption is justified by studies that demonstrated that peptides are 30 bound to HLA molecules and recognized by T cells in essentially an extended conformation. Derivation of specific algorithm coefficients has been described, for example, in Gulukota, K. *et al.*, *J. Mol. Biol.* 267:1258, 1997.

Additional methods to identify preferred peptide sequences, which also make use of specific motifs, include the use of neural networks and molecular modeling programs (see, e.g., Milik *et al.*, *Nature Biotechnology* 16:753, 1998; Altuvia *et al.*, *Hum. Immunol.* 58:1, 1997; Altuvia *et al.*, *J. Mol. Biol.* 249:244, 1995; Buus, S. *Curr. Opin. Immunol.* 11:209-213, 1999; Brusic, V. *et al.*, *Bioinformatics* 14:121-130, 1998; Parker *et al.*, *J. Immunol.* 152:163, 1993; Meister *et al.*, *Vaccine* 13:581, 1995; Hammer *et al.*, *J. Exp. Med.* 180:2353, 1994; Sturniolo *et al.*, *Nature Biotechnol.* 17:555 1999).

For example, it has been shown that in sets of A\*0201 motif-bearing peptides containing at least one preferred secondary anchor residue while avoiding the presence of any deleterious secondary anchor residues, 69% of the peptides will bind A\*0201 with an IC<sub>50</sub> less than 500 nM (Ruppert, J. *et al.* *Cell* 74:929, 1993). These algorithms are also flexible in that cut-off scores may be adjusted to select sets of peptides with greater or lower predicted binding properties, as desired.

In utilizing computer screening to identify peptide epitopes, a protein sequence or translated sequence may be analyzed using software developed to search for motifs, for example the "FINDPATTERNS" program (Devereux, *et al.* *Nucl. Acids Res.* 12:387-395, 1984) or MotifSearch 1.4 software program (D. Brown, San Diego, CA) to identify potential peptide sequences containing appropriate HLA binding motifs. The identified peptides can be scored using customized polynomial algorithms to predict their capacity to bind specific HLA class I or class II alleles. As appreciated by one of ordinary skill in the art, a large array of computer programming software and hardware options are available in the relevant art which can be employed to implement the motifs of the invention in order to evaluate (e.g., without limitation, to identify epitopes, identify epitope concentration per peptide length, or to generate analogs) known or unknown peptide sequences.

In accordance with the procedures described above, HCV peptide epitopes and analogs thereof that are able to bind HLA supertype groups or allele-specific HLA molecules have been identified (Tables VII-XX; Table XXII).

#### 30 IV.H. Preparation of Peptide Epitopes

Peptides in accordance with the invention can be prepared synthetically, by recombinant DNA technology or chemical synthesis, or from natural sources such as native tumors or pathogenic organisms. Peptide epitopes may be synthesized individually or as polyepitopic peptides. Although the peptide will preferably be substantially free of

other naturally occurring host cell proteins and fragments thereof, in some embodiments the peptides may be synthetically conjugated to native fragments or particles.

The peptides in accordance with the invention can be a variety of lengths, and either in their neutral (uncharged) forms or in forms which are salts. The peptides in 5 accordance with the invention are either free of modifications such as glycosylation, side chain oxidation, or phosphorylation; or they contain these modifications, subject to the condition that modifications do not destroy the biological activity of the peptides as described herein.

The peptides of the invention can be prepared in a wide variety of ways. For the 10 preferred relatively short size, the peptides can be synthesized in solution or on a solid support in accordance with conventional techniques. Various automatic synthesizers are commercially available and can be used in accordance with known protocols. (See, for example, Stewart & Young, SOLID PHASE PEPTIDE SYNTHESIS, 2D. ED., Pierce Chemical Co., 1984). Further, individual peptide epitopes can be joined using chemical ligation to 15 produce larger peptides that are still within the bounds of the invention.

Alternatively, recombinant DNA technology can be employed wherein a 20 nucleotide sequence which encodes an immunogenic peptide of interest is inserted into an expression vector, transformed or transfected into an appropriate host cell and cultivated under conditions suitable for expression. These procedures are generally known in the art, as described generally in Sambrook *et al.*, MOLECULAR CLONING, A LABORATORY 25 MANUAL, Cold Spring Harbor Press, Cold Spring Harbor, New York (1989). Thus, recombinant polypeptides which comprise one or more peptide sequences of the invention can be used to present the appropriate T cell epitope.

The nucleotide coding sequence for peptide epitopes of the preferred lengths 25 contemplated herein can be synthesized by chemical techniques, for example, the phosphotriester method of Matteucci, *et al.*, *J. Am. Chem. Soc.* 103:3185 (1981). Peptide analogs can be made simply by substituting the appropriate and desired nucleic acid base(s) for those that encode the native peptide sequence; exemplary nucleic acid substitutions are those that encode an amino acid defined by the motifs/supermotifs 30 herein. The coding sequence can then be provided with appropriate linkers and ligated into expression vectors commonly available in the art, and the vectors used to transform suitable hosts to produce the desired fusion protein. A number of such vectors and suitable host systems are now available. For expression of the fusion proteins, the coding sequence will be provided with operably linked start and stop codons, promoter and

terminator regions and usually a replication system to provide an expression vector for expression in the desired cellular host. For example, promoter sequences compatible with bacterial hosts are provided in plasmids containing convenient restriction sites for insertion of the desired coding sequence. The resulting expression vectors are 5 transformed into suitable bacterial hosts. Of course, yeast, insect or mammalian cell hosts may also be used, employing suitable vectors and control sequences.

It is often preferable that the peptide epitope be as small as possible while still maintaining substantially all of the immunologic activity of the native protein. When possible, it may be desirable to optimize HLA class I binding peptide epitopes of the 10 invention to a length of about 8 to about 13 amino acid residues, preferably 9 to 10. HLA class II binding peptide epitopes may be optimized to a length of about 6 to about 30 amino acids in length, preferably to between about 13 and about 20 residues. Preferably, the peptide epitopes are commensurate in size with endogenously processed pathogen-derived peptides or tumor cell peptides that are bound to the relevant HLA molecules, 15 however, the identification and preparation of peptides of other lengths can also be carried out using the techniques described herein.

In alternative embodiments, peptides of the invention can be linked as a polyepitopic peptide, or as a minigene that encodes a polyepitopic peptide.

In another embodiment, it is preferred to identify native peptide regions that 20 contain a high concentration of class I and/or class II epitopes. Such a sequence is generally selected on the basis that it contains the greatest number of epitopes per amino acid length. It is to be appreciated that epitopes can be present in a frame-shifted manner, *e.g.* a 10 amino acid long peptide could contain two 9 amino acid long epitopes and one 10 amino acid long epitope; upon intracellular processing, each epitope can be exposed 25 and bound by an HLA molecule upon administration of such a peptide. This larger, preferably multi-epitopic, peptide can be generated synthetically, recombinantly, or via cleavage from the native source.

#### IV.I. Assays to Detect T-Cell Responses

Once HLA binding peptides are identified, they can be tested for the ability to 30 elicit a T-cell response. The preparation and evaluation of motif-bearing peptides are described in PCT publications WO 94/20127 and WO 94/03205. Briefly, peptides comprising epitopes from a particular antigen are synthesized and tested for their ability to bind to the appropriate HLA proteins. These assays may involve evaluating the

binding of a peptide of the invention to purified HLA class I molecules in relation to the binding of a radioiodinated reference peptide. Alternatively, cells expressing empty class I molecules (*i.e.* lacking peptide therein) may be evaluated for peptide binding by immunofluorescent staining and flow microfluorimetry. Other assays that may be used to evaluate peptide binding include peptide-dependent class I assembly assays and/or the inhibition of CTL recognition by peptide competition. Those peptides that bind to the class I molecule, typically with an affinity of 500 nM or less, are further evaluated for their ability to serve as targets for CTLs derived from infected or immunized individuals, as well as for their capacity to induce primary *in vitro* or *in vivo* CTL responses that can give rise to CTL populations capable of reacting with selected target cells associated with a disease. Corresponding assays are used for evaluation of HLA class II binding peptides. HLA class II motif-bearing peptides that are shown to bind, typically at an affinity of 1000 nM or less, are further evaluated for the ability to stimulate HTL responses.

Conventional assays utilized to detect T cell responses include proliferation assays, lymphokine secretion assays, direct cytotoxicity assays, and limiting dilution assays. For example, antigen-presenting cells that have been incubated with a peptide can be assayed for the ability to induce CTL responses in responder cell populations. Antigen-presenting cells can be normal cells such as peripheral blood mononuclear cells or dendritic cells. Alternatively, mutant non-human mammalian cell lines that are deficient in their ability to load class I molecules with internally processed peptides and that have been transfected with the appropriate human class I gene, may be used to test for the capacity of the peptide to induce *in vitro* primary CTL responses.

Peripheral blood mononuclear cells (PBMCs) may be used as the responder cell source of CTL precursors. The appropriate antigen-presenting cells are incubated with peptide, after which the peptide-loaded antigen-presenting cells are then incubated with the responder cell population under optimized culture conditions. Positive CTL activation can be determined by assaying the culture for the presence of CTLs that kill radio-labeled target cells, both specific peptide-pulsed targets as well as target cells expressing endogenously processed forms of the antigen from which the peptide sequence was derived.

More recently, a method has been devised which allows direct quantification of antigen-specific T cells by staining with Fluorescein-labelled HLA tetrameric complexes (Altman, J. D. *et al.*, *Proc. Natl. Acad. Sci. USA* 90:10330, 1993; Altman, J. D. *et al.*, *Science* 274:94, 1996). Other relatively recent technical developments include staining

for intracellular lymphokines, and interferon release assays or ELISPOT assays.

Tetramer staining, intracellular lymphokine staining and ELISPOT assays all appear to be at least 10-fold more sensitive than more conventional assays (Lalvani, A. *et al.*, *J. Exp. Med.* 186:859, 1997; Dunbar, P. R. *et al.*, *Curr. Biol.* 8:413, 1998; Murali-Krishna, K. *et al.*, *Immunity* 8:177, 1998).

HTL activation may also be assessed using such techniques known to those in the art such as T cell proliferation and secretion of lymphokines, e.g. IL-2 (see, e.g. Alexander *et al.*, *Immunity* 1:751-761, 1994).

Alternatively, immunization of HLA transgenic mice can be used to determine immunogenicity of peptide epitopes. Several transgenic mouse models including mice with human A2.1, A11 (which can additionally be used to analyze HLA-A3 epitopes), and B7 alleles have been characterized and others (e.g., transgenic mice for HLA-A1 and A24) are being developed. HLA-DR1 and HLA-DR3 mouse models have also been developed. Additional transgenic mouse models with other HLA alleles may be generated as necessary. Mice may be immunized with peptides emulsified in Incomplete Freund's Adjuvant and the resulting T cells tested for their capacity to recognize peptide-pulsed target cells and target cells transfected with appropriate genes. CTL responses may be analyzed using cytotoxicity assays described above. Similarly, HTL responses may be analyzed using such assays as T cell proliferation or secretion of lymphokines.

Exemplary immunogenic peptide epitopes are set out in Table XXIII.

#### IV.J. Use of Peptide Epitopes as Diagnostic Agents and for Evaluating Immune Responses

In one embodiment of the invention, HLA class I and class II binding peptides as described herein can be used as reagents to evaluate an immune response. The immune response to be evaluated can be induced by using as an immunogen any agent that may result in the production of antigen-specific CTLs or HTLs that recognize and bind to the peptide epitope(s) to be employed as the reagent. The peptide reagent need not be used as the immunogen. Assay systems that can be used for such an analysis include relatively recent technical developments such as tetramers, staining for intracellular lymphokines and interferon release assays, or ELISPOT assays.

For example, a peptide of the invention may be used in a tetramer staining assay to assess peripheral blood mononuclear cells for the presence of antigen-specific CTLs following exposure to a tumor cell antigen or an immunogen. The HLA-tetrameric

complex is used to directly visualize antigen-specific CTLs (see, e.g., Ogg *et al.*, *Science* 279:2103-2106, 1998; and Altman *et al.*, *Science* 174:94-96, 1996) and determine the frequency of the antigen-specific CTL population in a sample of peripheral blood mononuclear cells. A tetramer reagent using a peptide of the invention may be generated 5 as follows: A peptide that binds to an HLA molecule is refolded in the presence of the corresponding HLA heavy chain and  $\beta_2$ -microglobulin to generate a trimolecular complex. The complex is biotinylated at the carboxyl terminal end of the heavy chain at a site that was previously engineered into the protein. Tetramer formation is then induced by the addition of streptavidin. By means of fluorescently labeled streptavidin, the 10 tetramer can be used to stain antigen-specific cells. The cells may then be identified, for example, by flow cytometry. Such an analysis may be used for diagnostic or prognostic purposes. Cells identified by the procedure can also be used for therapeutic purposes.

Peptides of the invention may also be used as reagents to evaluate immune recall responses. (see, e.g., Bertoni *et al.*, *J. Clin. Invest.* 100:503-513, 1997 and Penna *et al.*, *J. 15 Exp. Med.* 174:1565-1570, 1991.) For example, patient PBMC samples from individuals with HCV infection may be analyzed for the presence of antigen-specific CTLs or HTLs using specific peptides. A blood sample containing mononuclear cells may be evaluated by cultivating the PBMCs and stimulating the cells with a peptide of the invention. After an appropriate cultivation period, the expanded cell population may be analyzed, for 20 example, for cytotoxic activity (CTL) or for HTL activity.

The peptides may also be used as reagents to evaluate the efficacy of a vaccine. PBMCs obtained from a patient vaccinated with an immunogen may be analyzed using, for example, either of the methods described above. The patient is HLA typed, and peptide epitope reagents that recognize the allele-specific molecules present in that 25 patient are selected for the analysis. The immunogenicity of the vaccine is indicated by the presence of epitope-specific CTLs and/or HTLs in the PBMC sample.

The peptides of the invention may also be used to make antibodies, using techniques well known in the art (see, e.g. *CURRENT PROTOCOLS IN IMMUNOLOGY*, Wiley/Greene, NY; and *Antibodies A Laboratory Manual*, Harlow and Lane, Cold Spring 30 Harbor Laboratory Press, 1989), which may be useful as reagents to diagnose or monitor cancer. Such antibodies include those that recognize a peptide in the context of an HLA molecule, *i.e.*, antibodies that bind to a peptide-MHC complex.

#### IV.K. Vaccine Compositions

Vaccines and methods of preparing vaccines that contain an immunogenically effective amount of one or more peptides as described herein are further embodiments of the invention. Once appropriately immunogenic epitopes have been defined, they can be sorted and delivered by various means, herein referred to as "vaccine" compositions.

Such vaccine compositions can include, for example, lipopeptides (e.g., Vitiello, A. *et al.*, *J. Clin. Invest.* 95:341, 1995), peptide compositions encapsulated in poly(DL-lactide-co-

glycolide) ("PLG") microspheres (see, e.g., Eldridge, *et al.*, *Molec. Immunol.* 28:287-294, 1991; Alonso *et al.*, *Vaccine* 12:299-306, 1994; Jones *et al.*, *Vaccine* 13:675-681, 1995),

peptide compositions contained in immune stimulating complexes (ISCOMS) (see, e.g., Takahashi *et al.*, *Nature* 344:873-875, 1990; Hu *et al.*, *Clin Exp Immunol.* 113:235-243, 1998), multiple antigen peptide systems (MAPs) (see e.g., Tam, J. P., *Proc. Natl. Acad. Sci. U.S.A.* 85:5409-5413, 1988; Tam, J.P., *J. Immunol. Methods* 196:17-32, 1996), viral delivery vectors (Perkus, M. E. *et al.*, In: *Concepts in vaccine development*, Kaufmann, S.

H. E., ed., p. 379, 1996; Chakrabarti, S. *et al.*, *Nature* 320:535, 1986; Hu, S. L. *et al.*, *Nature* 320:537, 1986; Kieny, M.-P. *et al.*, *AIDS Bio/Technology* 4:790, 1986; Top, F. H. *et al.*, *J. Infect. Dis.* 124:148, 1971; Chanda, P. K. *et al.*, *Virology* 175:535, 1990), particles of viral or synthetic origin (e.g., Kofler, N. *et al.*, *J. Immunol. Methods*. 192:25, 1996; Eldridge, J. H. *et al.*, *Sem. Hematol.* 30:16, 1993; Falo, L. D., Jr. *et al.*, *Nature Med.* 7:649, 1995), adjuvants (Warren, H. S., Vogel, F. R., and Chedid, L. A. *Annu. Rev. Immunol.* 4:369, 1986; Gupta, R. K. *et al.*, *Vaccine* 11:293, 1993), liposomes (Reddy, R. *et al.*, *J. Immunol.* 148:1585, 1992; Rock, K. L., *Immunol. Today* 17:131, 1996), or,

naked or particle absorbed cDNA (Ulmer, J. B. *et al.*, *Science* 259:1745, 1993; Robinson, H. L., Hunt, L. A., and Webster, R. G., *Vaccine* 11:957, 1993; Shiver, J. W. *et al.*, In: *Concepts in vaccine development*, Kaufmann, S. H. E., ed., p. 423, 1996; Cease, K. B., and Berzofsky, J. A., *Annu. Rev. Immunol.* 12:923, 1994 and Eldridge, J. H. *et al.*, *Sem. Hematol.* 30:16, 1993). Toxin-targeted delivery technologies, also known as receptor mediated targeting, such as those of Avant Immunotherapeutics, Inc. (Needham, Massachusetts) may also be used.

Vaccines of the invention include nucleic acid-mediated modalities. DNA or RNA encoding one or more of the peptides of the invention can also be administered to a patient. This approach is described, for instance, in Wolff *et. al.*, *Science* 247:1465 (1990) as well as U.S. Patent Nos. 5,580,859; 5,589,466; 5,804,566; 5,739,118; 5,736,524; 5,679,647; WO 98/04720; and in more detail below. Examples of DNA-based

delivery technologies include "naked DNA", facilitated (bupivacaine, polymers, peptide-mediated) delivery, cationic lipid complexes, and particle-mediated ("gene gun") or pressure-mediated delivery (see, e.g., U.S. Patent No. 5,922,687).

For therapeutic or prophylactic immunization purposes, the peptides of the invention can also be expressed by viral or bacterial vectors. Examples of expression vectors include attenuated viral hosts, such as vaccinia or fowlpox. As an example of this approach, vaccinea virus is used as a vector to express nucleotide sequences that encode the peptides of the invention. Upon introduction into a host bearing a tumor, the recombinant vaccinia virus expresses the immunogenic peptide, and thereby elicits a host CTL and/or HTL response. Vaccinia vectors and methods useful in immunization protocols are described in, e.g., U.S. Patent No. 4,722,848. Another vector is BCG (Bacille Calmette Guerin). BCG vectors are described in Stover *et al.*, *Nature* 351:456-460 (1991). A wide variety of other vectors useful for therapeutic administration or immunization of the peptides of the invention, e.g. adeno and adeno-associated virus vectors, retroviral vectors, *Salmonella typhi* vectors, detoxified anthrax toxin vectors, and the like, will be apparent to those skilled in the art from the description herein.

Furthermore, vaccines in accordance with the invention encompass compositions of one or more of the claimed peptide(s). A peptide can be present in a vaccine individually. Alternatively, the peptide can exist as a homopolymer comprising multiple copies of the same peptide, or as a heteropolymer of various peptides. Polymers have the advantage of increased immunological reaction and, where different peptide epitopes are used to make up the polymer, the additional ability to induce antibodies and/or CTLs that react with different antigenic determinants of the pathogenic organism or tumor-related peptide targeted for an immune response. The composition can be a naturally occurring region of an antigen or can be prepared, e.g., recombinantly or by chemical synthesis.

Carriers that can be used with vaccines of the invention are well known in the art, and include, e.g., thyroglobulin, albumins such as human serum albumin, tetanus toxoid, polyamino acids such as poly L-lysine, poly L-glutamic acid, influenza, hepatitis B virus core protein, and the like. The vaccines can contain a physiologically tolerable (i.e., acceptable) diluent such as water, or saline, preferably phosphate buffered saline. The vaccines also typically include an adjuvant. Adjuvants such as incomplete Freund's adjuvant, aluminum phosphate, aluminum hydroxide, or alum are examples of materials well known in the art. Additionally, as disclosed herein, CTL responses can be primed by

conjugating peptides of the invention to lipids, such as tripalmitoyl-S-glycerylcysteinylseryl-serine (P<sub>3</sub>CSS).

Upon immunization with a peptide composition in accordance with the invention, via injection, aerosol, oral, transdermal, transmucosal, intrapleural, intrathecal, or other 5 suitable routes, the immune system of the host responds to the vaccine by producing large amounts of CTLs and/or HTLs specific for the desired antigen. Consequently, the host becomes at least partially immune to later infection, or at least partially resistant to developing an ongoing chronic infection, or derives at least some therapeutic benefit when the antigen was tumor-associated.

10 In some embodiments it may be desirable to combine the class I peptide components with components that induce or facilitate neutralizing antibody responses to the target antigen of interest, particularly to viral envelope antigens. A preferred embodiment of such a composition comprises class I and class II epitopes in accordance with the invention. An alternative embodiment of such a composition comprises a class I 15 and/or class II epitope in accordance with the invention, along with a PADRE™ (Epimmune, San Diego, CA) molecule (described, for example, in U.S. Patent Number 5,736,142).

20 A vaccine of the invention can also include antigen-presenting cells, such as dendritic cells, as a vehicle to present peptides of the invention. Vaccine compositions can be created *in vitro*, following dendritic cell mobilization and harvesting, whereby loading of dendritic cells occurs *in vitro*. For example, dendritic cells are transfected, e.g., with a minigene in accordance with the invention. The dendritic cell can then be administered to a patient to elicit immune responses *in vivo*.

25 Antigenic peptides are used to elicit a CTL and/or HTL response *ex vivo*, as well. The resulting CTL or HTL cells, can be used to treat tumors in patients that do not respond to other conventional forms of therapy, or will not respond to a therapeutic vaccine peptide or nucleic acid in accordance with the invention. *Ex vivo* CTL or HTL responses to a particular tumor-associated antigen are induced by incubating in tissue culture the patient's, or genetically compatible, CTL or HTL precursor cells together with 30 a source of antigen-presenting cells (APC), such as dendritic cells, and the appropriate immunogenic peptide. After an appropriate incubation time (typically about 7-28 days), in which the precursor cells are activated and expanded into effector cells, the cells are infused back into the patient, where they will destroy (CTL) or facilitate destruction

(HTL) of their specific target cell (an infected cell or a tumor cell). Transfected dendritic cells may also be used as antigen presenting cells.

The vaccine compositions of the invention can also be used in combination with antiviral drugs such as interferon- $\alpha$ , or other treatments for viral infection.

5 Preferably, the following principles are utilized when selecting an array of epitopes for inclusion in a polyepitopic composition for use in a vaccine, or for selecting discrete epitopes to be included in a vaccine and/or to be encoded by nucleic acids such as a minigene. It is preferred that each of the following principles are balanced in order to make the selection. The multiple epitopes to be incorporated in a given vaccine  
10 composition may be, but need not be, contiguous in sequence in the native antigen from which the epitopes are derived.

15 Preferably, the following principles are utilized when selecting an array of epitopes for inclusion in a polyepitopic composition for use in a vaccine, or for selecting discrete epitopes to be included in a vaccine and/or to be encoded by nucleic acids such as a minigene. Exemplary epitopes that may be utilized in a vaccine to treat or prevent HCV infection are set out in Tables XXVI-XXIX, and Table XXXII. It is preferred that each of the following principles are balanced in order to make the selection.

20 1.) Epitopes are selected which, upon administration, mimic immune responses that have been observed to be correlated with HCV clearance. For HLA Class I this includes 3-4 epitopes that come from at least one antigen of HCV. For HLA Class II a similar rationale is employed; again 3-4 epitopes are selected from at least one HCV antigen (see e.g., Rosenberg *et al.*, *Science* 278:1447-1450).

25 2.) Epitopes are selected that have the requisite binding affinity established to be correlated with immunogenicity: for HLA Class I an  $IC_{50}$  of 500 nM or less, or for Class II an  $IC_{50}$  of 1000 nM or less.

30 3.) Sufficient supermotif bearing-peptides, or a sufficient array of allele-specific motif-bearing peptides, are selected to give broad population coverage. For example, it is preferable to have at least 80% population coverage. A Monte Carlo analysis, a statistical evaluation known in the art, can be employed to assess the breadth, or redundancy of, population coverage.

35 4.) When selecting epitopes from cancer-related antigens it is often preferred to select analogs because the patient may have developed tolerance to the native epitope.

When selecting epitopes for infectious disease-related antigens it is preferable to select either native or analoged epitopes.

5.) Of particular relevance are epitopes referred to as "nested epitopes."

Nested epitopes occur where at least two epitopes overlap in a given peptide sequence. A

5 nested peptide sequence can comprise both HLA class I and HLA class II epitopes.

When providing nested epitopes, it is preferable to provide a sequence that has the greatest number of epitopes per provided sequence. Preferably, one avoids providing a peptide that is any longer than the amino terminus of the amino terminal epitope and the carboxyl terminus of the carboxyl terminal epitope in the peptide. When providing a

10 longer peptide sequence, such as a sequence comprising nested epitopes, it is important to screen the sequence in order to insure that it does not have pathological or other deleterious biological properties.

6.) If a polyepitopic protein is created, or when creating a minigene, an objective is to generate the smallest peptide that encompasses the epitopes of interest.

15 This principle is similar, if not the same as that employed when selecting a peptide comprising nested epitopes. However, with an artificial polyepitopic peptide, the size minimization objective is balanced against the need to integrate any spacer sequences between epitopes in the polyepitopic protein. Spacer amino acid residues can be introduced to avoid junctional epitopes (an epitope recognized by the immune system, not

20 present in the target antigen, and only created by the man-made juxtaposition of epitopes), or to facilitate cleavage between epitopes and thereby enhance epitope presentation. Junctional epitopes are generally to be avoided because the recipient may

generate an immune response to that non-native epitope. Of particular concern is a junctional epitope that is a "dominant epitope." A dominant epitope may lead to such a

25 zealous response that immune responses to other epitopes are diminished or suppressed.

Examples of polyepitopic vaccine compositions designed based on the above criteria can include epitopes from the core, S, E1, NS1/E2, NS2, NS3, NS4, and NS5 domains of the HCV polyprotein. These regions encompass the following amino acid sequences using numbering relative to the prototype HCV-1 strain (Genbank accession

30 number M62321; *see, e.g.*, US Patent Nos. 5,683,864 and 5,670,153): C domain (amino acids 1-120); S (amino acids 120-400); NS3 (amino acids 1050-1640); NS4 (amino acids 1640-2000); NS5 (amino acids 2000-3011); and envelop proteins, E1 and E2/NS1, encompassing amino acids 192-750. Amino acids 750 to 1050 are designated as domain X as applied to the present invention. As appreciated by one of ordinary skill in the art,

the designation of the amino acid range for each domain may diverge to some extent from that of HCV-1 depending on the strain of HCV. One of ordinary skill in the art, when looking at an HCV polyprotein sequence, would readily be able to determine the domain boundaries.

- 5        Specific embodiments of the polyepitopic compositions of the present invention include a pharmaceutical composition comprising a pharmaceutically acceptable carrier and combination of motif-bearing peptides that are immunologically cross-reactive with peptides of HCV-1, wherein at least one of the peptides bears a motif of Table Ia, and further wherein the combination of motif-bearing peptides consists of: a) one or more
- 10      peptides comprising at least 8 amino acids from an HCV C domain; b) one or more peptides comprising at least 8 amino acids of a further domain selected from the group consisting of: an S domain, an NS3 domain, an NS4 domain, or an NS5 domain, and; c) optionally, one or more motif-bearing peptides from one or more additional HCV domains with a *proviso* that an additional domain is not a further domain listed in "b".
- 15      Preferably, such a pharmaceutical composition may additionally comprise one or more distinct HCV motif-bearing peptide(s) comprising at least 8 amino acids of an X domain or, alternatively, the composition may further comprise additional HCV motif-bearing peptide(s) that are from an envelope domain, the envelope domain peptide(s) consisting of one or more copies of a single HCV peptide comprising at least 8 amino acids of an
- 20      envelope domain.

In another embodiment, the polyepitopic pharmaceutical composition may comprise a pharmaceutically acceptable carrier and combination of motif-bearing peptides that are immunologically cross-reactive with HCV-1 peptides, the peptides from multiple domains of HCV, wherein at least one of the peptides bears a motif of Table Ia, and wherein the combination of motif-bearing peptides consists essentially of: a) one or more peptides comprising at least 8 amino acids from a C domain; and, b) one or more peptides comprising at least 8 amino acids from an S, NS3, NS4, or NS5 domain, and, one HCV peptide comprising at least 8 amino acids of an envelope domain. Such a composition may further comprise one or more HCV motif-bearing peptides comprising at least 8 amino acids of an X domain.

Alternatively, a pharmaceutical composition of the invention may comprise: a) a pharmaceutically acceptable carrier; and, b) a combination of one or more motif-bearing peptides of at least 8 amino acids derived from one or more hepatitis C virus (HCV) domains, wherein said peptides are cross-reactive with peptides of HCV-1, with a *proviso*

that the combination does not include a peptide of at least 8 amino acids from an HCV C domain, and wherein at least one of the peptides bears a motif of Table Ia, said domains selected from the group consisting of: an S domain; an NS3 domain; an NS4 domain; an NS5 domain; and, an X domain. Such a composition may additionally comprise motif-bearing HCV envelope peptide(s) consisting of one or more copies of a single HCV peptide comprising at least 8 amino acids of an envelope domain.

Lastly, an embodiment of the invention may comprise a pharmaceutical composition comprising a pharmaceutically acceptable carrier and combination of two or more motif-bearing peptides from a single domain of an HCV-1 strain, said peptides 10 immunologically cross-reactive with peptides of an HCV-1 antigen, wherein at least one of the peptides bears a motif of Table Ia, and the peptides are derived from HCV, and the HCV domain is selected from the group consisting of: a C domain; an S domain; an NS3 domain; an NS4 domain; an NS5 domain; an X domain; or, an envelope domain from a single HCV strain, with a *proviso* that the envelope domain is other than a variable 15 envelope domain.

In the embodiments set forth, "peptides immunologically cross-reactive with HCV-1" refers to peptides that are bound by the same antibody; "derived from" refers to a fragment or subsequence and conservatively modified variants thereof.

#### 20 IV.K.1. Minigene Vaccines

A number of different approaches are available which allow simultaneous delivery of multiple epitopes. Nucleic acids encoding the peptides of the invention are a particularly useful embodiment of the invention. Epitopes for inclusion in a minigene are preferably selected according to the guidelines set forth in the previous section. A 25 preferred means of administering nucleic acids encoding the peptides of the invention uses minigene constructs encoding a peptide comprising one or multiple epitopes of the invention.

The use of multi-epitope minigenes is described below and in, e.g., co-pending application U.S.S.N. 09/311,784; An, L. and Whitton, J. L., *J. Virol.* 71:2292, 1997; 30 Thomson, S. A. *et al.*, *J. Immunol.* 157:822, 1996; Whitton, J. L. *et al.*, *J. Virol.* 67:348, 1993; Hanke, R. *et al.*, *Vaccine* 16:426, 1998. For example, a multi-epitope DNA plasmid encoding supermotif- and/or motif-bearing HCV epitopes derived from multiple regions of the HCV polyprotein sequence, the PADRE™ universal helper T cell epitope (or

multiple HTL epitopes from HCV), and an endoplasmic reticulum-translocating signal sequence can be engineered.

The immunogenicity of a multi-epitopic minigene can be tested in transgenic mice to evaluate the magnitude of CTL induction responses against the epitopes tested.

5 Further, the immunogenicity of DNA-encoded epitopes *in vivo* can be correlated with the *in vitro* responses of specific CTL lines against target cells transfected with the DNA plasmid. Thus, these experiments can show that the minigene serves to both: 1.) generate a CTL response and 2.) that the induced CTLs recognized cells expressing the encoded epitopes.

10 For example, to create a DNA sequence encoding the selected epitopes (minigene) for expression in human cells, the amino acid sequences of the epitopes may be reverse translated. A human codon usage table can be used to guide the codon choice for each amino acid. These epitope-encoding DNA sequences may be directly adjoined, so that when translated, a continuous polypeptide sequence is created. To optimize expression 15 and/or immunogenicity, additional elements can be incorporated into the minigene design. Examples of amino acid sequences that can be reverse translated and included in the minigene sequence include: HLA class I epitopes, HLA class II epitopes, a ubiquitination signal sequence, and/or an endoplasmic reticulum targeting signal. In addition, HLA presentation of CTL and HTL epitopes may be improved by including 20 synthetic (*e.g.* poly-alanine) or naturally-occurring flanking sequences adjacent to the CTL or HTL epitopes; these larger peptides comprising the epitope(s) are within the scope of the invention.

The minigene sequence may be converted to DNA by assembling oligonucleotides that encode the plus and minus strands of the minigene. Overlapping oligonucleotides 25 (30-100 bases long) may be synthesized, phosphorylated, purified and annealed under appropriate conditions using well known techniques. The ends of the oligonucleotides can be joined, for example, using T4 DNA ligase. This synthetic minigene, encoding the epitope polypeptide, can then be cloned into a desired expression vector.

Standard regulatory sequences well known to those of skill in the art are 30 preferably included in the vector to ensure expression in the target cells. Several vector elements are desirable: a promoter with a down-stream cloning site for minigene insertion; a polyadenylation signal for efficient transcription termination; an *E. coli* origin of replication; and an *E. coli* selectable marker (*e.g.* ampicillin or kanamycin resistance). Numerous promoters can be used for this purpose, *e.g.*, the human cytomegalovirus

(hCMV) promoter. See, e.g., U.S. Patent Nos. 5,580,859 and 5,589,466 for other suitable promoter sequences.

Additional vector modifications may be desired to optimize minigene expression and immunogenicity. In some cases, introns are required for efficient gene expression, 5 and one or more synthetic or naturally-occurring introns could be incorporated into the transcribed region of the minigene. The inclusion of mRNA stabilization sequences and sequences for replication in mammalian cells may also be considered for increasing minigene expression.

Once an expression vector is selected, the minigene is cloned into the polylinker 10 region downstream of the promoter. This plasmid is transformed into an appropriate *E. coli* strain, and DNA is prepared using standard techniques. The orientation and DNA sequence of the minigene, as well as all other elements included in the vector, are confirmed using restriction mapping and DNA sequence analysis. Bacterial cells harboring the correct plasmid can be stored as a master cell bank and a working cell bank.

15 In addition, immunostimulatory sequences (ISSs or CpGs) appear to play a role in the immunogenicity of DNA vaccines. These sequences may be included in the vector, outside the minigene coding sequence, if desired to enhance immunogenicity.

In some embodiments, a bi-cistronic expression vector which allows production of 20 both the minigene-encoded epitopes and a second protein (included to enhance or decrease immunogenicity) can be used. Examples of proteins or polypeptides that could beneficially enhance the immune response if co-expressed include cytokines (e.g., IL-2, IL-12, GM-CSF), cytokine-inducing molecules (e.g., LeIF), costimulatory molecules, or for HTL responses, pan-DR binding proteins (PADRE<sup>TM</sup>, Epimmune, San Diego, CA). Helper (HTL) epitopes can be joined to intracellular targeting signals and expressed 25 separately from expressed CTL epitopes; this allows direction of the HTL epitopes to a cell compartment different than that of the CTL epitopes. If required, this could facilitate more efficient entry of HTL epitopes into the HLA class II pathway, thereby improving HTL induction. In contrast to HTL or CTL induction, specifically decreasing the immune response by co-expression of immunosuppressive molecules (e.g. TGF- $\beta$ ) may be 30 beneficial in certain diseases.

Therapeutic quantities of plasmid DNA can be produced for example, by fermentation in *E. coli*, followed by purification. Aliquots from the working cell bank are used to inoculate growth medium, and grown to saturation in shaker flasks or a bioreactor

according to well known techniques. Plasmid DNA can be purified using standard bioseparation technologies such as solid phase anion-exchange resins supplied by QIAGEN, Inc. (Valencia, California). If required, supercoiled DNA can be isolated from the open circular and linear forms using gel electrophoresis or other methods.

5 Purified plasmid DNA can be prepared for injection using a variety of formulations. The simplest of these is reconstitution of lyophilized DNA in sterile phosphate-buffer saline (PBS). This approach, known as "naked DNA," is currently being used for intramuscular (IM) administration in clinical trials. To maximize the immunotherapeutic effects of minigene DNA vaccines, an alternative method for  
10 formulating purified plasmid DNA may be desirable. A variety of methods have been described, and new techniques may become available. Cationic lipids can also be used in the formulation (see, *e.g.*, as described by WO 93/24640; Mannino & Gould-Fogerite, *BioTechniques* 6(7): 682 (1988); U.S. Pat No. 5,279,833; WO 91/06309; and Felgner, *et al.*, *Proc. Nat'l Acad. Sci. USA* 84:7413 (1987). In addition, glycolipids, fusogenic  
15 liposomes, peptides and compounds referred to collectively as protective, interactive, non-condensing compounds (PINC) could also be complexed to purified plasmid DNA to influence variables such as stability, intramuscular dispersion, or trafficking to specific organs or cell types.

Target cell sensitization can be used as a functional assay for expression and HLA  
20 class I presentation of minigene-encoded CTL epitopes. For example, the plasmid DNA is introduced into a mammalian cell line that is suitable as a target for standard CTL chromium release assays. The transfection method used will be dependent on the final formulation. Electroporation can be used for "naked" DNA, whereas cationic lipids allow direct *in vitro* transfection. A plasmid expressing green fluorescent protein (GFP) can be  
25 co-transfected to allow enrichment of transfected cells using fluorescence activated cell sorting (FACS). These cells are then chromium-51 ( $^{51}\text{Cr}$ ) labeled and used as target cells for epitope-specific CTL lines; cytolysis, detected by  $^{51}\text{Cr}$  release, indicates both production of, and HLA presentation of, minigene-encoded CTL epitopes. Expression of HLA epitopes may be evaluated in an analogous manner using assays to assess HLA  
30 activity.

*In vivo* immunogenicity is a second approach for functional testing of minigene DNA formulations. Transgenic mice expressing appropriate human HLA proteins are immunized with the DNA product. The dose and route of administration are formulation dependent (*e.g.*, IM for DNA in PBS, intraperitoneal (IP) for lipid-complexed DNA).

Twenty-one days after immunization, splenocytes are harvested and restimulated for 1 week in the presence of peptides encoding each epitope being tested. Thereafter, for CTL effector cells, assays are conducted for cytolysis of peptide-loaded, <sup>51</sup>Cr-labeled target cells using standard techniques. Lysis of target cells that were sensitized by HLA loaded 5 with peptide epitopes, corresponding to minigene-encoded epitopes, demonstrates DNA vaccine function for *in vivo* induction of CTLs. Immunogenicity of HTL epitopes is evaluated in transgenic mice in an analogous manner.

Alternatively, the nucleic acids can be administered using ballistic delivery as described, for instance, in U.S. Patent No. 5,204,253. Using this technique, particles 10 comprised solely of DNA are administered. In a further alternative embodiment, DNA can be adhered to particles, such as gold particles.

#### **IV.K.2. Combinations of CTL Peptides with Helper Peptides**

Vaccine compositions comprising the peptides of the present invention, or analogs 15 thereof, which have immunostimulatory activity may be modified to provide desired attributes, such as improved serum half life, or to enhance immunogenicity.

For instance, the ability of the peptides to induce CTL activity can be enhanced by linking the peptide to a sequence which contains at least one epitope that is capable of inducing a T helper cell response. The use of T helper epitopes in conjunction with CTL 20 epitopes to enhance immunogenicity is illustrated, for example, in co-pending U.S.S.N. 08/820360, U.S.S.N. 08/197,484, and U.S.S.N. 08/464,234.

Particularly preferred CTL epitope/HTL epitope conjugates are linked by a spacer molecule. The spacer is typically comprised of relatively small, neutral molecules, such as amino acids or amino acid mimetics, which are substantially uncharged under 25 physiological conditions. The spacers are typically selected from, *e.g.*, Ala, Gly, or other neutral spacers of nonpolar amino acids or neutral polar amino acids. It will be understood that the optionally present spacer need not be comprised of the same residues and thus may be a hetero- or homo-oligomer. When present, the spacer will usually be at least one or two residues, more usually three to six residues. Alternatively, the CTL 30 peptide may be linked to the T helper peptide without a spacer.

Although the CTL peptide epitope can be linked directly to the T helper peptide epitope, often CTL epitope/HTL epitope conjugates are linked by a spacer molecule. The spacer is typically comprised of relatively small, neutral molecules, such as amino acids or amino acid mimetics, which are substantially uncharged under physiological

conditions. The spacers are typically selected from, *e.g.*, Ala, Gly, or other neutral spacers of nonpolar amino acids or neutral polar amino acids. It will be understood that the optionally present spacer need not be comprised of the same residues and thus may be a hetero- or homo-oligomer. When present, the spacer will usually be at least one or two residues, more usually three to six residues. The CTL peptide epitope can be linked to the T helper peptide epitope either directly or via a spacer either at the amino or carboxy terminus of the CTL peptide. The amino terminus of either the immunogenic peptide or the T helper peptide may be acylated.

HTL peptide epitopes can also be modified to alter their biological properties. For example, peptides comprising HTL epitopes can contain D-amino acids to increase their resistance to proteases and thus extend their serum half-life. Also, the epitope peptides of the invention can be conjugated to other molecules such as lipids, proteins or sugars, or any other synthetic compounds, to increase their biological activity. Specifically, the T helper peptide can be conjugated to one or more palmitic acid chains at either the amino or carboxyl termini.

In certain embodiments, the T helper peptide is one that is recognized by T helper cells present in the majority of the population. This can be accomplished by selecting amino acid sequences that bind to many, most, or all of the HLA class II molecules. These are known as "loosely HLA-restricted" or "promiscuous" T helper sequences.

Examples of amino acid sequences that are promiscuous include sequences from antigens such as tetanus toxoid at positions 830-843 (QYIKANSKFIGITE), *Plasmodium falciparum* CS protein at positions 378-398 (DIEKKIAKMEKASSVFNVVNS), and *Streptococcus* 18kD protein at positions 116 (GAVDSILGGVATYGAA). Other examples include peptides bearing a DR 1-4-7 supermotif, or either of the DR3 motifs.

Alternatively, it is possible to prepare synthetic peptides capable of stimulating T helper lymphocytes, in a loosely HLA-restricted fashion, using amino acid sequences not found in nature (*see, e.g.*, PCT publication WO 95/07707). These synthetic compounds called Pan-DR-binding epitopes (*e.g.*, PADRE™, Epimmune, Inc., San Diego, CA) are designed to most preferably bind most HLA-DR (human HLA class II) molecules. For instance, a pan-DR-binding epitope peptide having the formula: aKXVVWANTLKAa, where "X" is either cyclohexylalanine, phenylalanine, or tyrosine, and a is either D-alanine or L-alanine, has been found to bind to most HLA-DR alleles, and to stimulate the response of T helper lymphocytes from most individuals, regardless of their HLA type.

An alternative of a pan-DR binding epitope comprises all "L" natural amino acids and can be provided in the form of nucleic acids that encode the epitope.

In some embodiments it may be desirable to include in the pharmaceutical compositions of the invention at least one component which primes cytotoxic T

5 lymphocytes. Lipids have been identified as agents capable of priming CTL *in vivo* against viral antigens. For example, palmitic acid residues can be attached to the ε-and α-amino groups of a lysine residue and then linked, *e.g.*, via one or more linking residues such as Gly, Gly-Gly-, Ser, Ser-Ser, or the like, to an immunogenic peptide. The lipidated peptide can then be administered either directly in a micelle or particle, 10 incorporated into a liposome, or emulsified in an adjuvant, *e.g.*, incomplete Freund's adjuvant. In a preferred embodiment, a particularly effective immunogenic comprises palmitic acid attached to ε- and α- amino groups of Lys, which is attached via linkage, *e.g.*, Ser-Ser, to the amino terminus of the immunogenic peptide.

As another example of lipid priming of CTL responses, *E. coli* lipoproteins, such 15 as tripalmitoyl-S-glycercylcysteinylseryl- serine (P<sub>3</sub>CSS) can be used to prime virus specific CTL when covalently attached to an appropriate peptide. (See, *e.g.*, Deres, *et al.*, *Nature* 342:561, 1989). Peptides of the invention can be coupled to P<sub>3</sub>CSS, for example, and the lipopeptide administered to an individual to specifically prime a CTL response to the target antigen. Moreover, because the induction of neutralizing antibodies can also be 20 primed with P<sub>3</sub>CSS-conjugated epitopes, two such compositions can be combined to more effectively elicit both humoral and cell-mediated responses to infection.

As noted herein, additional amino acids can be added to the termini of a peptide to provide for ease of linking peptides one to another, for coupling to a carrier support or larger peptide, for modifying the physical or chemical properties of the peptide or 25 oligopeptide, or the like. Amino acids such as tyrosine, cysteine, lysine, glutamic or aspartic acid, or the like, can be introduced at the C- or N-terminus of the peptide or oligopeptide, particularly class I peptides. However, it is to be noted that modification at the carboxyl terminus of a CTL epitope may, in some cases, alter binding characteristics 30 of the peptide. In addition, the peptide or oligopeptide sequences can differ from the natural sequence by being modified by terminal-NH<sub>2</sub> acylation, *e.g.*, by alkanoyl (C<sub>1</sub>-C<sub>20</sub>) or thioglycolyl acetylation, terminal-carboxyl amidation, *e.g.*, ammonia, methylamine, *etc.* In some instances these modifications may provide sites for linking to a support or other molecule.

*Vaccine Compositions Comprising Dendritic Cells Pulsed with CTL and/or HTL Peptides*

An embodiment of a vaccine composition in accordance with the invention comprises *ex vivo* administration of a cocktail of epitope-bearing peptides to PBMC, or 5 isolated DC therefrom, from the patient's blood. A pharmaceutical to facilitate harvesting of DC can be used, such as GM-CSF/IL-4. After pulsing the DC with peptides and prior to reinfusion into patients, the DC are washed to remove unbound peptides. In this embodiment, a vaccine comprises peptide-pulsed DCs which present the pulsed peptide epitopes complexed with HLA molecules on their surfaces. The vaccine is then 10 administered to the patient.

**IV.L. Administration of Vaccines for Therapeutic or Prophylactic Purposes**

The peptides of the present invention and pharmaceutical and vaccine compositions of the invention are useful for administration to mammals, particularly 15 humans, to treat and/or prevent HCV infection. Vaccine compositions containing the peptides of the invention are administered to a patient infected with HCV or to an individual susceptible to, or otherwise at risk for, HCV infection to elicit an immune response against HCV antigens and thus enhance the patient's own immune response capabilities. In therapeutic applications, peptide and/or nucleic acid compositions are 20 administered to a patient in an amount sufficient to elicit an effective CTL and/or HTL response to the virus antigen and to cure or at least partially arrest or slow symptoms and/or complications. An amount adequate to accomplish this is defined as "therapeutically effective dose." Amounts effective for this use will depend on, *e.g.*, the particular composition administered, the manner of administration, the stage and severity 25 of the disease being treated, the weight and general state of health of the patient, and the judgment of the prescribing physician.

The vaccine compositions of the invention may also be used purely as prophylactic agents. Generally the dosage for an initial prophylactic immunization 30 generally occurs in a unit dosage range where the lower value is about 1, 5, 50, 500, or 1000  $\mu$ g and the higher value is about 10,000; 20,000; 30,000; or 50,000  $\mu$ g. Dosage values for a human typically range from about 500  $\mu$ g to about 50,000  $\mu$ g per 70 kilogram patient. This is followed by boosting dosages of between about 1.0  $\mu$ g to about 50,000  $\mu$ g of peptide administered at defined intervals from about four weeks to six months after the

initial administration of vaccine. The immunogenicity of the vaccine may be assessed by measuring the specific activity of CTL and HTL obtained from a sample of the patient's blood.

As noted above, peptides comprising CTL and/or HTL epitopes of the invention 5 induce immune responses when presented by HLA molecules and contacted with a CTL or HTL specific for an epitope comprised by the peptide. The manner in which the peptide is contacted with the CTL or HTL is not critical to the invention. For instance, the peptide can be contacted with the CTL or HTL either *in vivo* or *in vitro*. If the contacting occurs *in vivo*, the peptide itself can be administered to the patient, or other 10 vehicles, *e.g.*, DNA vectors encoding one or more peptides, viral vectors encoding the peptide(s), liposomes and the like, can be used, as described herein. When the peptide is contacted *in vitro*, the vaccinating agent can comprise a population of cells, *e.g.*, peptide-pulsed dendritic cells, or TAA-specific CTLs, which have been induced by pulsing antigen-presenting cells *in vitro* with the peptide. Such a cell population is subsequently 15 administered to a patient in a therapeutically effective dose.

The peptides or DNA encoding them can be administered individually or as fusions of one or more peptide sequences.

For pharmaceutical compositions, the immunogenic peptides of the invention, or 20 DNA encoding them, are generally administered to an individual already infected with HCV. The peptides or DNA encoding them can be administered individually or as fusions of one or more peptide sequences. Those in the incubation phase or the acute phase of infection can be treated with the immunogenic peptides separately or in conjunction with other treatments, as appropriate.

For therapeutic use, administration should generally begin at the first diagnosis of 25 HCV infection. This is followed by boosting doses until at least symptoms are substantially abated and for a period thereafter. In chronic infection, loading doses followed by boosting doses may be required.

Treatment of an infected individual with the compositions of the invention may 30 hasten resolution of the infection in acutely infected individuals. For those individuals susceptible (or predisposed) to developing chronic infection, the compositions are particularly useful in methods for preventing the evolution from acute to chronic infection. Where susceptible individuals are identified prior to or during infection, the composition can be targeted to them, thus minimizing the need for administration to a larger population.

The peptide or other compositions used for the treatment or prophylaxis of HCV infection can be used, *e.g.*, in persons who have not manifested symptoms of disease but who act as a disease vector. In this context, it is generally important to provide an amount of the peptide epitope delivered by a mode of administration sufficient to

5 effectively stimulate a cytotoxic T cell response; compositions which stimulate helper T cell responses can also be given in accordance with this embodiment of the invention.

The dosage for an initial therapeutic immunization generally occurs in a unit dosage range where the lower value is about 1, 5, 50, 500, or 1000  $\mu$ g and the higher value is about 10,000; 20,000; 30,000; or 50,000  $\mu$ g. Dosage values for a human

10 typically range from about 500  $\mu$ g to about 50,000  $\mu$ g per 70 kilogram patient. Boosting dosages of between about 1.0  $\mu$ g to about 50000  $\mu$ g of peptide pursuant to a boosting regimen over weeks to months may be administered depending upon the patient's response and condition as determined by measuring the specific activity of CTL and HTL obtained from the patient's blood. The peptides and compositions of the present

15 invention may be employed in serious disease states, that is, life-threatening or potentially life threatening situations. In such cases, as a result of the minimal amounts of extraneous substances and the relative nontoxic nature of the peptides in preferred compositions of the invention, it is possible and may be felt desirable by the treating physician to administer substantial excesses of these peptide compositions relative to

20 these stated dosage amounts.

Thus, for treatment of chronic infection, a representative dose is in the range disclosed above, namely where the lower value is about 1, 5, 50, 500, or 1000  $\mu$ g and the higher value is about 10,000; 20,000; 30,000; or 50,000  $\mu$ g, preferably from about 500  $\mu$ g to about 50,000  $\mu$ g per 70 kilogram patient. Initial doses followed by boosting doses at

25 established intervals, *e.g.*, from four weeks to six months, may be required, possibly for a prolonged period of time to effectively immunize an individual. In the case of chronic infection, administration should continue until at least clinical symptoms or laboratory tests indicate that the viral infection has been eliminated or substantially abated and for a period thereafter. The dosages, routes of administration, and dose schedules are adjusted

30 in accordance with methodologies known in the art.

The pharmaceutical compositions for therapeutic treatment are intended for parenteral, topical, oral, intrathecal, or local administration. Preferably, the pharmaceutical compositions are administered parentally, *e.g.*, intravenously,

subcutaneously, intradermally, or intramuscularly. Thus, the invention provides compositions for parenteral administration which comprise a solution of the immunogenic peptides dissolved or suspended in an acceptable carrier, preferably an aqueous carrier. A variety of aqueous carriers may be used, *e.g.*, water, buffered water, 0.8% saline, 0.3% 5 glycine, hyaluronic acid and the like. These compositions may be sterilized by conventional, well known sterilization techniques, or may be sterile filtered. The resulting aqueous solutions may be packaged for use as is, or lyophilized, the lyophilized preparation being combined with a sterile solution prior to administration. The compositions may contain pharmaceutically acceptable auxiliary substances as required 10 to approximate physiological conditions, such as pH-adjusting and buffering agents, tonicity adjusting agents, wetting agents, preservatives, and the like, for example, sodium acetate, sodium lactate, sodium chloride, potassium chloride, calcium chloride, sorbitan monolaurate, triethanolamine oleate, *etc.*

The concentration of peptides of the invention in the pharmaceutical formulations 15 can vary widely, *i.e.*, from less than about 0.1%, usually at or at least about 2% to as much as 20% to 50% or more by weight, and will be selected primarily by fluid volumes, viscosities, *etc.*, in accordance with the particular mode of administration selected.

A human unit dose form of the peptide composition is typically included in a pharmaceutical composition that comprises a human unit dose of an acceptable carrier, 20 preferably an aqueous carrier, and is administered in a volume of fluid that is known by those of skill in the art to be used for administration of such compositions to humans (*see, e.g.*, Remington's Pharmaceutical Sciences, 17<sup>th</sup> Edition, A. Gennaro, Editor, Mack Publishing Co., Easton, Pennsylvania, 1985).

The peptides of the invention may also be administered via liposomes, which 25 serve to target the peptides to a particular tissue, such as lymphoid tissue, or to target selectively to infected cells, as well as to increase the half-life of the peptide composition. Liposomes include emulsions, foams, micelles, insoluble monolayers, liquid crystals, phospholipid dispersions, lamellar layers and the like. In these preparations, the peptide to be delivered is incorporated as part of a liposome, alone or in conjunction with a 30 molecule which binds to a receptor prevalent among lymphoid cells, such as monoclonal antibodies which bind to the CD45 antigen, or with other therapeutic or immunogenic compositions. Thus, liposomes either filled or decorated with a desired peptide of the invention can be directed to the site of lymphoid cells, where the liposomes then deliver the peptide compositions. Liposomes for use in accordance with the invention are formed

from standard vesicle-forming lipids, which generally include neutral and negatively charged phospholipids and a sterol, such as cholesterol. The selection of lipids is generally guided by consideration of, *e.g.*, liposome size, acid lability and stability of the liposomes in the blood stream. A variety of methods are available for preparing 5 liposomes, as described in, *e.g.*, Szoka, *et al.*, *Ann. Rev. Biophys. Bioeng.* 9:467 (1980), and U.S. Patent Nos. 4,235,871, 4,501,728, 4,837,028, and 5,019,369.

For targeting cells of the immune system, a ligand to be incorporated into the liposome can include, *e.g.*, antibodies or fragments thereof specific for cell surface determinants of the desired immune system cells. A liposome suspension containing a 10 peptide may be administered intravenously, locally, topically, *etc.* in a dose which varies according to, *inter alia*, the manner of administration, the peptide being delivered, and the stage of the disease being treated.

For solid compositions, conventional nontoxic solid carriers may be used which include, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium 15 stearate, sodium saccharin, talcum, cellulose, glucose, sucrose, magnesium carbonate, and the like. For oral administration, a pharmaceutically acceptable nontoxic composition is formed by incorporating any of the normally employed excipients, such as those carriers previously listed, and generally 10-95% of active ingredient, that is, one or more peptides of the invention, and more preferably at a concentration of 25%-75%.

20 For aerosol administration, the immunogenic peptides are preferably supplied in finely divided form along with a surfactant and propellant. Typical percentages of peptides are 0.01%-20% by weight, preferably 1%-10%. The surfactant must, of course, be nontoxic, and preferably soluble in the propellant. Representative of such agents are the esters or partial esters of fatty acids containing from 6 to 22 carbon atoms, such as 25 caproic, octanoic, lauric, palmitic, stearic, linoleic, linolenic, olesteric and oleic acids with an aliphatic polyhydric alcohol or its cyclic anhydride. Mixed esters, such as mixed or natural glycerides may be employed. The surfactant may constitute 0.1%-20% by weight of the composition, preferably 0.25-5%. The balance of the composition is ordinarily propellant. A carrier can also be included, as desired, as with, *e.g.*, lecithin for intranasal 30 delivery.

#### IV.M. Kits

The peptide and nucleic acid compositions of this invention can be provided in kit form together with instructions for vaccine administration. Typically the kit would

include desired peptide compositions in a container, preferably in unit dosage form and instructions for administration. An alternative kit would include a minigene construct with desired nucleic acids of the invention in a container, preferably in unit dosage form together with instructions for administration. Lymphokines such as IL-2 or IL-12 may 5 also be included in the kit. Other kit components that may also be desirable include, for example, a sterile syringe, booster dosages, and other desired excipients.

The invention will be described in greater detail by way of specific examples. The following examples are offered for illustrative purposes, and are not intended to limit 10 the invention in any manner. Those of skill in the art will readily recognize a variety of non-critical parameters that can be changed or modified to yield alternative embodiments in accordance with the invention.

## V. EXAMPLES

As in many viral diseases, there is evidence that clearance of HCV is mediated by 15 CTL. In a study of primary HCV infection in six chimpanzees, four progressed to chronic infection (Cooper *et al.*, abstract, 19<sup>th</sup> US-Japan Hepatitis Joint Panel Meeting, January 27-29, 1998). It was found that these four animals showed either no CTL response or a very narrowly focused response during early infection. In contrast, in the 20 remaining two animals that resolved the infection, a broad CTL response was observed against multiple HCV proteins, some of which were conserved. Weiner *et al.* (*Proc. Natl. Acad. Sci. USA* 92:2755-2759, 1995) demonstrated that viral escape, in which the epitopes presented to PATR class I molecules mutated, was linked with a progression 25 toward chronic infection. These data show a role for the CTL in directing the course of HCV disease, and in shaping the genetic composition of HCV species in the persistently infected host.

In work in humans, Koziel and co-workers have established the presence of HCV-specific CTL in liver infiltrates from patients with chronic HCV infection (Koziel *et al.*, *J. Immunol.* 149:3339, 1992; and Koziel *et al.*, *J. Virol.* 67:7522, 1993), and have also identified a number of CTL epitopes recognized in the context of several different HLA 30 class I molecules. Other investigators have shown that HCV-specific CTL can be detected in the peripheral blood of patients with chronic hepatitis C (Cerny *et al.*, *J. Clin. Invest.* 95:521, 1995; Cerny *et al.*, *Curr. Topics in Micro. and Immunol.* 189:169, 1994; Cerny *et al.*, Abst. 2<sup>nd</sup> International Meeting on Hepatitis C and Related Viruses; La Jolla, CA, 1994; Battegay *et al.*, Abst. 2<sup>nd</sup> International Meeting on Hepatitis C and Related

Viruses; La Jolla, CA, 1994; Shirai *et al.*, *J. Virol.* 68:3334, 1994; Shirai *et al.*, *J. Immunol.* 154:2733, 1995; Battegay *et al.*, *J. Virol.* 69:2462, 1995). In addition, escape variants have been demonstrated in patients chronically infected with HCV (Chang *et al.*, *J. Clin. Invest.* 100:2376-2385, 1997; Tsai *et al.*, *Gastroenterology* 115:954-966, 1998).

5 The magnitude of the CTL responses observed in HCV-infected patients is, in general, higher than those observed in the case of chronic hepatitis B infection, suggesting that there is less impairment of specific T cell immunity than with HBV infection. The magnitude of CTL responses in HCV patients is, however, lower than those observed in HBV infected individuals who successfully cleared HBV infection.

10 These results support the understanding that HCV infected patients are capable of responding to active immunotherapy, and suggest that potentiation and increasing of T cell responses to HCV may be of use in therapy and prevention of chronic HCV infection (Prince, A. M. *FEMS Micro. Rev.* 14:273, 1994).

Several groups have analyzed the potential role of HCV-specific CTL responses in disease resistance and pathogenesis. In some studies no correlation was found between CTL viremia and CTL precursor frequency for individual HCV epitopes (Rehermann *et al.*, *J. Clin. Invest.* 98:1432-1440, 1996; Wong *et al.*, *J. Immunol.* 160:1479-1488, 1998). In other studies, however, it was shown that a clear correlation existed between levels of HCV infection and CTL responses, provided that the global response against multiple 20 CTL epitopes was considered (Rehermann *et al.*, *J. Virol.* 70:7092-7102, 1996). These data represent a strong rationale for development of vaccine constructs capable of inducing vigorous CTL responses directed against a multiplicity of conserved HCV-derived epitopes.

Koziel and colleagues have demonstrated the presence of HCV-specific CTLs, as 25 well as T helper cell responses, in exposed but seronegative individuals (Koziel *et al.*, *J. Infect. Diseases* 176:859-866, 1997). In addition, HCV-specific CTLs have been detected in healthy, seronegative family members of chronically HCV-infected patients, indicating that a protective immunity is established in absence of a detectable infection (Bronowicki *et al.*, *J. Infect. Dis.* 176:518-522, 1997; Scognamiglio *et al.*, in preparation).

30 Experimental evidence also indicates that HTL epitopes play an important role in immune reactivity and defenses against HCV infection (Missale *et al.*, *J. Clin. Invest.* 98:706-714, 1996). Diepolder *et al.* (in *Lancet* 346:1006, 1995) have shown that a region of the NS3 gene (NS3 1007-1534) is recognized by patients who clear acute HCV infection, but is not seen by patients who develop chronic infection. Subsequent studies

showed that this particular region contain a highly cross-reactive HTL epitope (NS3 1248-1261), which binds with good affinity to 10 of 13 DR molecules tested, and is highly conserved in 30/33 different HCV isolates considered (Diepolder *et al.*, *J. Virol.* 71:6011-6019, 1997). These data suggested that directing HTL responses to this type of epitope (rather than to less cross-reactive and/or highly variable ones) will be of therapeutic and prophylactic benefit and strongly argue for inclusion of this and other epitopes with similar characteristics in HCV vaccine constructs.

5 The following examples illustrate identification, selection, and use of immunogenic Class I and Class II peptide epitopes for inclusion in vaccine compositions.

10

Example 1: HLA Class I and Class II Binding Assays

The following example of peptide binding to HLA molecules demonstrates quantification of binding affinities of HLA class I and class II peptides. Binding assays can be performed with peptides that are either motif-bearing or not motif-bearing.

15 Epstein-Barr virus (EBV)-transformed homozygous cell lines, fibroblasts, CIR, or 721.22 transfectants were used as sources of HLA class I molecules. The specific cell lines routinely used for purification of MHC class I and class II molecules are listed in Table XXIV. Cell lysates were prepared and HLA molecules purified in accordance with disclosed protocols (Sidney *et al.*, *Current Protocols in Immunology* 18.3.1 (1998);  
20 Sidney, *et al.*, *J. Immunol.* 154:247 (1995); Sette, *et al.*, *Mol. Immunol.* 31:813 (1994)). HLA molecules were purified from lysates by affinity chromatography. The lysate was passed over a column of Sepharose CL-4B beads coupled to an appropriate antibody. The antibodies used for the extraction of HLA from cell lysates are listed in Table XXV. The anti-HLA column was then washed with 10mM Tris-HCL, pH 8.0, in 1% NP-40,  
25 PBS, and PBS containing 0.4% n-octylglucoside and HLA molecules were eluted with 50mM diethylamine in 0.15M NaCl containing 0.4% n-octylglucoside, pH 11.5. A 1/25 volume of 2.0M Tris, pH 6.8, was added to the eluate to reduce the pH to ~8.0. Eluates were then be concentrated by centrifugation in Centriprep 30 concentrators (Amicon, Beverly, MA). Protein content was evaluated by a BCA protein assay (Pierce Chemical  
30 Co., Rockford, IL) and confirmed by SDS-PAGE.

A detailed description of the protocol utilized to measure the binding of peptides to Class I and Class II MHC has been published (Sette *et al.*, *Mol. Immunol.* 31:813, 1994; Sidney *et al.*, in *Current Protocols in Immunology*, Margulies, Ed., John Wiley & Sons, New York, Section 18.3, 1998). Briefly, purified MHC molecules (5 to 500nM)

were incubated with various unlabeled peptide inhibitors and 1-10nM  $^{125}\text{I}$ -radiolabeled probe peptides for 48h in PBS containing 0.05% Nonidet P-40 (NP40) (or 20% w/v digitonin for H-2 IA assays) in the presence of a protease inhibitor cocktail. All assays were at pH 7.0 with the exception of DRB1\*0301, which was performed at pH 4.5, and 5 DRB1\*1601 (DR2w21 $\beta_1$ ) and DRB4\*0101 (DRw53), which were performed at pH 5.0.

Following incubation, MHC-peptide complexes were separated from free peptide by gel filtration on 7.8 mm x 15 cm TSK200 columns (TosoHaas 16215, Montgomeryville, PA). Because the large size of the radiolabeled peptide used for the DRB1\*1501 (DR2w2 $\beta_1$ ) assay makes separation of bound from unbound peaks more 10 difficult under these conditions, all DRB1\*1501 (DR2w2 $\beta_1$ ) assays were performed using a 7.8mm x 30cm TSK2000 column eluted at 0.6 mls/min. The eluate from the TSK columns was passed through a Beckman 170 radioisotope detector, and radioactivity was plotted and integrated using a Hewlett-Packard 3396A integrator, and the fraction of peptide bound was determined.

15 Radiolabeled peptides were iodinated using the chloramine-T method. Representative radiolabeled probe peptides utilized in each assay, and its assay specific IC<sub>50</sub> nM, are summarized in Tables IV and V. Typically, in preliminary experiments, each MHC preparation was titered in the presence of fixed amounts of radiolabeled peptides to determine the concentration of HLA molecules necessary to bind 10-20% of 20 the total radioactivity. All subsequent inhibition and direct binding assays were performed using these HLA concentrations.

Since under these conditions [label]<[HLA] and IC<sub>50</sub>≥[HLA], the measured IC<sub>50</sub> values are reasonable approximations of the true K<sub>D</sub> values. Peptide inhibitors are typically tested at concentrations ranging from 120  $\mu\text{g}/\text{ml}$  to 1.2 ng/ml, and are tested in 25 two to four completely independent experiments. To allow comparison of the data obtained in different experiments, a relative binding figure is calculated for each peptide by dividing the IC<sub>50</sub> of a positive control for inhibition by the IC<sub>50</sub> for each tested peptide (typically unlabeled versions of the radiolabeled probe peptide). For database purposes, and inter-experiment comparisons, relative binding values are compiled. These values 30 can subsequently be converted back into IC<sub>50</sub> nM values by dividing the IC<sub>50</sub> nM of the positive controls for inhibition by the relative binding of the peptide of interest. This method of data compilation has proven to be the most accurate and consistent for

comparing peptides that have been tested on different days, or with different lots of purified MHC.

Because the antibody used for HLA-DR purification (LB3.1) is  $\alpha$ -chain specific,  $\beta_1$  molecules are not separated from  $\beta_3$  (and/or  $\beta_4$  and  $\beta_5$ ) molecules. The  $\beta_1$  specificity of the binding assay is obvious in the cases of DRB1\*0101 (DR1), DRB1\*0802 (DR8w2), and DRB1\*0803 (DR8w3), where no  $\beta_3$  is expressed. It has also been demonstrated for DRB1\*0301 (DR3) and DRB3\*0101 (DR52a), DRB1\*0401 (DR4w4), DRB1\*0404 (DR4w14), DRB1\*0405 (DR4w15), DRB1\*1101 (DR5), DRB1\*1201 (DR5w12), DRB1\*1302 (DR6w19) and DRB1\*0701 (DR7). The problem of  $\beta$  chain specificity for DRB1\*1501 (DR2w2 $\beta_1$ ), DRB5\*0101 (DR2w2 $\beta_2$ ), DRB1\*1601 (DR2w21 $\beta_1$ ), DRB5\*0201 (DR51Dw21), and DRB4\*0101 (DRw53) assays is circumvented by the use of fibroblasts. Development and validation of assays with regard to DR $\beta$  molecule specificity have been described previously (see, e.g., Southwood *et al.*, *J. Immunol.* 160:3363-3373, 1998).

Binding assays as outlined above may be used to analyze supermotif and/or motif-bearing epitopes as, for example, described in Example 2.

Example 2. Identification of Conserved HLA Supermotif- and Motif-Bearing CTL Candidate Epitopes

Vaccine compositions of the invention may include multiple epitopes that comprise multiple HLA supermotifs or motifs to achieve broad population coverage. This example illustrates the identification of supermotif- and motif-bearing epitopes for the inclusion in such a vaccine composition. Calculation of population coverage was performed using the strategy described below.

*Computer searches and algorithms for identification of supermotif and/or motif-bearing epitopes*

Computer searches for epitopes bearing HLA Class I or Class II supermotifs or motifs were performed as follows. All translated HCV isolate sequences were analyzed using a text string search software program, *e.g.*, MotifSearch 1.4 (D. Brown, San Diego) to identify potential peptide sequences containing appropriate HLA binding motifs; alternative programs are readily produced in accordance with information in the art in view of the motif/supermotif disclosure herein. Furthermore, such calculations can be

made mentally. Identified A2-, A3-, and DR-supermotif sequences were scored using polynomial algorithms to predict their capacity to bind to specific HLA-Class I or Class II molecules. These polynomial algorithms take into account both extended and refined motifs (that is, to account for the impact of different amino acids at different positions), 5 and are essentially based on the premise that the overall affinity (or  $\Delta G$ ) of peptide-HLA molecule interactions can be approximated as a linear polynomial function of the type:

$$\Delta G = a_{1i} \times a_{2i} \times a_{3i} \dots \times a_{ni}$$

where  $a_{ji}$  is a coefficient which represents the effect of the presence of a given amino acid (j) at a given position (i) along the sequence of a peptide of n amino acids. The crucial 10 assumption of this method is that the effects at each position are essentially independent of each other (i.e., independent binding of individual side-chains). When residue  $j$  occurs at position  $i$  in the peptide, it is assumed to contribute a constant amount  $j_i$  to the free energy of binding of the peptide irrespective of the sequence of the rest of the peptide. This assumption is justified by studies from our laboratories that demonstrated that 15 peptides are bound to MHC and recognized by T cells in essentially an extended conformation (data omitted herein).

The method of derivation of specific algorithm coefficients has been described in Gulukota *et al.*, *J. Mol. Biol.* 267:1258-126, 1997; (see also Sidney *et al.*, *Human Immunol.* 45:79-93, 1996; and Southwood *et al.*, *J. Immunol.* 160:3363-3373, 1998). 20 Briefly, for all  $i$  positions, anchor and non-anchor alike, the geometric mean of the average relative binding (ARB) of all peptides carrying  $j$  is calculated relative to the remainder of the group, and used as the estimate of  $j_i$ . For Class II peptides, if multiple alignments are possible, only the highest scoring alignment is utilized, following an iterative procedure. To calculate an algorithm score of a given peptide in a test set, the 25 ARB values corresponding to the sequence of the peptide are multiplied. If this product exceeds a chosen threshold, the peptide is predicted to bind. Appropriate thresholds are chosen as a function of the degree of stringency of prediction desired.

#### *Selection of HLA-A2 supertype cross-reactive peptides*

30 Complete polyprotein sequences from fourteen HCV isolates were aligned, then scanned, utilizing motif identification software, to identify conserved 9- and 10-mer sequences containing the HLA-A2-supermotif main anchor specificity.

A total of 231 conserved, HLA-A2 supermotif-positive sequences were identified. These peptides were then evaluated for the presence of A\*0201 preferred secondary anchor residues using A\*0201-specific polynomial algorithms. A total of 67 conserved, motif-bearing and algorithm-positive sequences were identified.

5       Fifty of these conserved, motif-containing 9- and 10-mer peptides were tested for their capacity to bind to purified HLA-A\*0201 molecules *in vitro* (HLA-A\*0201 is considered a prototype A2 supertype molecule). Sixteen peptides bound A\*0201 with IC<sub>50</sub> values ≤500 nM; 4 with high binding affinities (IC<sub>50</sub> values ≤50 nM) and 12 with intermediate binding affinities, in the 50-500 nM range (Table XXVI).

10      These 16 peptides were then tested for binding to additional A2-supertype molecules (A\*0202, A\*0203, A\*0206, and A\*6802). As shown in Table XXVI, most of these peptides were found to be A2-supertype cross-reactive binders. More specifically, 12/16 (75%) peptides bound at least three of the five A2-supertype molecules tested.

15      *Selection of HLA-A3 supermotif-bearing epitopes*

The sequences from the same fourteen known HCV isolates scanned above were also examined for the presence of conserved peptides with the HLA-A3-supermotif primary anchors. A total of 71 conserved 9- or 10-mer motif containing sequences were identified. Further analysis using the A03 and A11 algorithms (see, e.g., Gulukota *et al.*, 20 *J. Mol. Biol.* 267:1258-1267, 1997 and Sidney *et al.*, *Human Immunol.* 45:79-93, 1996) identified 39 sequences that scored high in either or both algorithms. Twenty seven of the 39 peptides were synthesized and tested for binding to HLA-A\*03 and HLA-A\*11, the two most prevalent A3-supertype molecules. Fifteen peptides were identified which bound A3 and/or A11 with binding affinities of ≤500 nM (Table XXVII). These peptides were then tested for binding cross-reactivity to the other common A3-supertype alleles (A\*3101, A\*3301, and A\*6801). Seven of the 15 peptides bound at least three of the five HLA-A3-supertype molecules tested.

In the course of an independent series of experiments (Kubo *et al.*, *J. Immunol.* 152:3913-3924, 1994), one peptide, HCV NS3 1262, not identified by the selection criteria utilized above because it does not have the A3-supermotif main anchor specificity, was determined to be cross-reactive in the A3-supertype, binding A\*03, A\*11, and A\*6801. It is also shown in Table XXVII. Interestingly, this peptide

represents a single residue N-terminal truncation of peptide 1073.14, which is also shown in Table XXVII.

In summary, 8 peptides that bind 3 or more A3-supertype molecules derived from conserved regions of the HCV genome were identified.

5

*Selection of HLA-B7 supermotif bearing epitopes*

When the same fourteen HCV isolates were also analyzed for the presence of conserved 9- or 10-mer peptides with the HLA-B7-supermotif, 35 sequences were identified. The corresponding peptides were synthesized and tested for binding to HLA-

10 B\*0702, the most common B7-supertype allele (*i.e.*, the prototype B7 supertype allele).

Thirteen peptides bound B\*0702 with IC<sub>50</sub> of  $\leq$ 500 nM (Table XXVIIIa). These 13 peptides were then tested for binding to other common B7-supertype molecules (B\*3501, B\*51, B\*5301, and B\*5401). As shown in Table XXVIIIa, only 1 peptide (Core 169) was capable of binding to three or more of the five B7-supertype alleles tested.

15 To identify additional B7-supertype epitopes, further studies were undertaken.

The protein sequences from the fourteen HCV isolates utilized above were again examined to identify conserved, motif-containing 8- and 11-mers. The isolates were also examined for 9- and 10-mer sequences allowing for lower conservancy (51%-78%).

20 Twenty-five 8-mers, sixteen 11-mers, and thirty-five 9- and 10-mers were identified, synthesized, and tested for binding to B\*0702. Thirteen peptides bound with high or intermediate affinity (IC<sub>50</sub>  $\leq$ 500 nM) (Table XXVIIIb). These peptides were additionally screened for binding to other B7-supertype molecules. Only one cross-reactive binder, the NS3 1378 8-mer (peptide 29.0035/1260.04), was identified (Table XXVIIIb).

25 In summary, a total of two cross-reactive B7-supertype binders were identified (Core 169 and NS3 1378).

*Selection of A1 and A24 motif-bearing epitopes*

To further increase population coverage, HLA-A1 and -A24 epitopes can also be incorporated into potential vaccine constructs.

30 In a previous analysis, two A1 and three A24 binders, 100% conserved among four strains of HCV, were identified (Wentworth *et al.*, *Int. Immunol.* 8:651-659, 1996). An analysis of the protein sequence data from the fourteen HCV strains utilized above demonstrated that these peptides were >79% conserved, and also identified an additional

eleven A1- and twenty five A24-motif-containing conserved sequences (see Table XXIXA and B). Eight of the additional eleven A1 peptides and seven of the additional twenty five A24 peptides were tested for binding to the appropriate HLA molecule (i.e., A1 or A24). Overall, as shown in Table XXIX, four A1-motif peptides (A) and three 5 A24-motif peptides (B) have been found with binding capacities of 500 nM or less for the appropriate allele-specific HLA molecule.

Analysis of the HLA-A2 and A3 supermotif-bearing epitopes identified above revealed that in 13/14 cases, peptides binding the supertype prototype HLA molecule (i.e. A\*0201 for the A2 supertype, and A\*0301 for the A3 supertype) with an IC<sub>50</sub> of less than 10 100nM were cross-reactive and recognized by HCV-infected patients as described in Example 3, which follows. Based on these observations, two A1 peptides and one A24 peptide epitopes were also selected as candidates for inclusion in vaccine compositions; these peptides bind the appropriate HLA molecule with an IC<sub>50</sub> of less than 100nM.

15 Example 3: Confirmation of Immunogenicity

*Evaluation of A\*0201 immunogenicity*

It has been shown that CTL induced in A\*0201/K<sup>b</sup> transgenic mice exhibit specificity similar to CTL induced in the human system (see, e.g., Vitiello *et al.*, *J. Exp. Med.* 173:1007-1015, 1991; Wentworth *et al.*, *Eur. J. Immunol.* 26:97-101, 1996).

20 Accordingly, these mice were used to evaluate the immunogenicity of the twelve conserved A2-supertype cross-reactive peptides identified in Example 2 above.

CTL induction in transgenic mice following peptide immunization has been described (Vitiello *et al.*, *J. Exp. Med.* 173:1007-1015, 1991; Alexander *et al.*; *J. Immunol.* 159:4753-4761, 1997). In these studies, mice were injected subcutaneously at 25 the base of the tail with each peptide (50 µg/mouse) emulsified in IFA in the presence of an excess of an IA<sup>b</sup>-restricted helper peptide (140 µg/mouse) (HBV core 128-140, Sette *et al.*, *J. Immunol.* 153:5586-5592, 1994). Eleven days after injection, splenocytes were incubated in the presence of peptide-loaded syngenic LPS blasts. After six days, cultures were assayed for cytotoxic activity using peptide-pulsed targets. The data, summarized in 30 Table XXX, indicate that 7 of the 12 peptides (58%) were capable of inducing primary CTL responses in A\*0201/K<sup>b</sup> transgenic mice. (For these studies, a peptide was considered positive if it induced CTL (L.U. 30/10<sup>6</sup> cells ≥2 in at least two transgenic animals (Wentworth *et al.*, *Eur. J. Immunol.* 26:97-101, 1996).

The conserved, cross reactive candidate CTL epitopes were also tested for recognition *in vitro* by PBMCs obtained from HCV-infected patients. Briefly, PBMC from patients infected with HCV were cultured in the presence of 10 µg/ml of synthetic peptide. After 7 and 14 days, the cultures were restimulated with peptide. The cultures 5 were assayed for cytolytic activity on day 21 using target cells pulsed with the specific peptide in a standard four hour <sup>51</sup>Cr release assay. The data are summarized in Table XXX. As shown, all 12 peptides are CTL epitopes recognized by PBMC from HCV-infected patients. From the data in Table XXX, it is interesting to note that HLA 10 transgenics did not fully reveal the immunogenicity of some peptides that were positive in recall responses. This apparent discrepancy may reflect differences in the route of immunization utilized (e.g., natural infection versus peptide immunization), or CTL repertoire.

*Evaluation of A\*03/A11 immunogenicity*

15 The immunogenicity of six of the eight A3-supertype cross-reactive peptides identified in Example 2 above was evaluated in HLA-A11/K<sup>b</sup> transgenic mice, using the protocol described above for HLA-A2 transgenic mice (Alexander *et al.*, *J. Immunol.* 159:4753-4761, 1997). Five of these six peptides were able to induce primary CTL 159:4753-4761, 1997). Five of these six peptides were able to induce primary CTL responses (Table XXXI).

20 All eight peptides were also studied by collaborators using PBMC cultures from HCV infected patients and contacts of such patients. This data is also summarized in Table XXXI. Briefly, all eight peptides were recognized by HCV infected individuals.

*Evaluation of B7 immunogenicity*

25 One of the two B7-supertype cross-reactive peptides (1145.12, Core 169) has been evaluated for immunogenicity in HCV-infected patients. Two independent collaborators have shown that this peptide is indeed immunogenic, and is recognized by T cells from HCV-infected patients (Chang *et al.*, *J. Immunol.* 162:1156-1164, 1999)

30 Example 4: Implementation of the Extended Supermotif to Improve the Binding Capacity of Native Epitopes by Creating Analogs

HLA motifs and supermotifs (comprising primary and/or secondary residues) are useful in the identification and preparation of highly cross-reactive native peptides, as demonstrated herein. Moreover, the definition of HLA motifs and supermotifs also

allows one to engineer highly cross-reactive epitopes by identifying residues within a native peptide sequence which can be analogued, or "fixed" to confer upon the peptide certain characteristics, *e.g.* greater cross-reactivity within the group of HLA molecules that comprise a supertype, and/or greater binding affinity for some or all of those HLA 5 molecules. Examples of analog peptides that exhibit modulated binding affinity are set forth in this example.

#### *Analoging at Primary Anchor Residues*

As shown in Example 2, more than ten different HCV-derived, A2-supertype-restricted epitopes were identified. Peptide engineering strategies are implemented to further increase the cross-reactivity of the candidate epitopes identified above which bind 10 3/5 of the A2 supertype alleles tested. On the basis of the data disclosed, *e.g.*, in related and co-pending U.S.S.N 09/226,775, the main anchors of A2-supermotif-bearing peptides are altered, for example, to introduce a preferred L, I, V, or M at position 2, and I or V at 15 the C-terminus.

To analyze the cross-reactivity of the analog peptides, each engineered analog is initially tested for binding to the prototype A2 supertype allele A\*0201, then, if A\*0201 binding capacity is maintained, for A2-supertype cross-reactivity.

Similarly, analogs of HLA-A3 supermotif-bearing epitopes may also be 20 generated. For example, peptides binding to 3/5 of the A3-supertype molecules may be engineered at primary anchor residues to possess a preferred residue (V, S, M, or A) at position 2.

The analog peptides are then tested for the ability to bind A\*03 and A\*11 (prototype A3 supertype alleles). Those peptides that demonstrate  $\leq$  500 nM binding 25 capacity are then tested for A3-supertype cross-reactivity.

Similarly to the A2- and A3- motif bearing peptides, peptides binding 3 or more B7-supertype alleles may be improved, where possible, to achieve increased cross-reactive binding. B7 supermotif-bearing peptides may, for example, be engineered to possess a preferred residue (V, I, L, or F) at the C-terminal primary anchor position, as 30 demonstrated by Sidney *et al.* (*J. Immunol.* 157:3480-3490, 1996).

#### *Analoging at Secondary Anchor Residues*

Moreover, HLA supermotifs are of value in engineering highly cross-reactive peptides and/or peptides that bind HLA molecules with increased affinity by identifying

particular residues at secondary anchor positions that are associated with such properties. Demonstrating this, the binding capacity of a peptide representing a discreet single amino acid substitution at position one was analyzed. Peptide 1145.13 (Table XXVIIIC), which represents the substitution of L to F at position 1 of the core 169 sequence, binds all five 5 B7-supertype molecules with a good affinity (all  $IC_{50}$  values  $\leq 132$  nM), and in 3 instances has higher affinity over that of the parent peptide by >35-fold.

Because so few B7-supertype cross-reactive epitopes were identified, our results from previous binding evaluations were analyzed to identify conserved (8-, 9-, 10-, or 11-mer) peptides which bind, minimally, 3/5 B7 supertype molecules with weak affinity 10 ( $IC_{50}$  of 500nM-5 $\mu$ M). This analysis identified 9 peptides, 6 of which are analogued (including core 169 which had been previously analogued). These peptides are tested for enhanced binding affinity and B7-supertype cross-reactivity.

Engineered analogs with sufficiently improved binding capacity or cross-reactivity are tested for immunogenicity in HLA-B7-transgenic mice, following for 15 example, IFA immunization or lipopeptide immunization.

In conclusion, these data demonstrate that by the use of even single amino acid substitutions, it is possible to increase the binding affinity and/or cross-reactivity of peptide ligands for HLA supertype molecules.

20 Example 5: Identification of conserved HCV-derived sequences with HLA-DR binding motifs

Peptide epitopes bearing an HLA class II supermotif or motif may also be identified as outlined below using methodology similar to that described in Examples 1-3.

25 *Selection of HLA-DR-supermotif-bearing epitopes*

To identify HCV-derived, HLA class II HTL epitopes, the same fourteen HCV polyprotein sequences used for the identification of HLA Class I supermotif/motif sequences were analyzed for the presence of sequences bearing an HLA-DR-motif or supermotif. Specifically, 15-mer sequences were selected comprising a DR-supermotif, 30 further comprising a 9-mer core, and three-residue N- and C-terminal flanking regions (15 amino acids total). It was also required that the 15-mer sequence be conserved in at least 79% (11/14) of the HCV strains analyzed. These criteria identified a total of 49 non-redundant sequences, which are shown in Table XXXIIA. (In the context of Class II

epitopes, a sequence is considered operationally redundant if more than 80% of it's sequence overlaps with another peptide.)

Protocols for predicting peptide binding to DR molecules have been developed (Southwood *et al.*, *J. Immunol.* 160:3363-3373, 1998). These protocols, specific for 5 individual DR molecules, allow the scoring, and ranking, of 9-mer core regions. Each protocol not only scores peptide sequences for the presence of DR-supermotif primary anchors (i.e., at position 1 and position 6) within a 9-mer core, but additionally evaluates sequences for the presence of secondary anchors. Using allele specific selection tables (see, *e.g.*, Southwood *et al.*, *ibid.*), it has been found that these protocols efficiently select 10 peptide sequences with a high probability of binding a particular DR molecule.

Additionally, it has been found that performing these protocols in tandem, specifically those for DR1, DR4w4, and DR7, can efficiently select DR cross-reactive peptides.

To see if these protocols serve to identify additional epitopes, the same HCV polyproteins used above were re-scanned for the presence of 15-mer peptides with 9-mer 15 core regions that were  $\geq 79\%$  (11/14 strains) conserved. This identified 152 sequences; 49 of which were identified previously, as described above. Next, the 9-mer core region of each of these peptides was scored using the DR1, DR4w4, and DR7 algorithms. Twenty-two peptides, including 12 new sequences (10 peptides were from the original set of 49) 20 were found to have 9-mer cores with protocol-derived scores predictive of cross-reactive DR binders. The 12 additional sequences are shown in Table XXXIIB.

The conserved, HCV-derived peptides identified above were tested for their binding capacity for various common HLA-DR molecules. All peptides were initially tested for binding to the DR molecules in the primary panel: DR1, DR4w4, and DR7. Peptides binding at least 2 of these 3 DR molecules were then tested for binding to 25 DR2w2  $\beta$ 1, DR2w2  $\beta$ 2, DR6w19, and DR9 molecules in secondary assays. Finally, peptides binding at least 2 of the 4 secondary panel DR molecules, and thus cumulatively at least 4 of 7 different DR molecules, were screened for binding to DR4w15, DR5w11, and DR8w2 molecules in tertiary assays. Peptides binding at least 7 of the 10 DR molecules comprising the primary, secondary, and tertiary screening assays were 30 considered cross-reactive DR binders. The composition of these screening panels, and the phenotypic frequency of associated antigens, are shown in Table XXXIII.

Upon testing, it was found that 29 of the original 75 peptides (39%) bound two or more of the primary HLA molecules. Twenty-six of these cross-reactive binders were

then tested in the secondary assays, and nineteen were found to bind at least four of the seven HLA DR molecules in the primary and secondary panels. Finally, the nineteen peptides passing the secondary screening phase were tested for binding in the tertiary assays. As a result, nine peptides were identified which bound at least seven of ten common HLA-DR molecules. Table XXXIV shows these nine peptides and their binding capacity for each allele-specific HLA-DR molecule in the primary through tertiary panels. Also shown in Table XXXIV are two peptides (F134.05 and F134.08) for which a complete binding analysis was not performed. However, both of these peptides bound six of the seven HLA DR molecules tested. F134.08 nests peptide 1283.44, which bound eight of 10 allele-specific HLA molecules.

In conclusion, eleven cross-reactive DR-binding peptides, derived from six discrete (*i.e.* non-redundant) regions of the HCV genome, have been identified. Two of the six regions from which these epitopes were derived are covered by multiple, overlapping epitopes.

15

#### *Selection of conserved DR3 motif peptides*

Because HLA-DR3 is an allele that is prevalent in Caucasian, Black, and Hispanic populations, DR3 binding capacity is an important criterion in the selection of HLA epitopes. However, data generated previously indicated that DR3 only rarely cross-reacts with other DR alleles (Sidney *et al.*, *J. Immunol.* 149:2634-2640, 1992; Geluk *et al.*, *J. Immunol.* 152:5742-5748, 1994; Southwood *et al.*, *J. Immunol.* 160:3363-3373, 1998). This is not entirely surprising in that the DR3 peptide-binding motif appears to be distinct from the specificity of most other DR alleles.

To efficiently identify peptides that bind DR3, target proteins were analyzed for conserved sequences carrying one of the two DR3 specific binding motifs reported by Geluk *et al.* (*J. Immunol.* 152:5742-5748, 1994). Fifteen sequences, including a peptide nested within a DR-supermotif sequence identified above (peptide Pape 22), were identified (Table XXXId). Preferably, DR3 motifs will be found clustered in proximity with DR supermotif regions.

30 Fourteen of the fifteen peptides containing a DR3 motif were tested for their DR3 binding capacity. Two peptides (CH35.0106 and CH35.0107) were found to bind DR3 with an affinity of 1 $\mu$ M or less (Table XXXV), and thereby qualify as HLA class II high affinity binders.

DR3 binding epitopes identified in this manner may then be included in vaccine compositions with DR supermotif-bearing peptide epitopes.

Example 6: Immunogenicity of candidate HCV-derived HTL epitopes and known dominant HCV HTL epitope

In the course of collaborative studies with G. Pape and C. Ferrari, eight conserved, HCV-derived peptides have been identified which are recognized by HCV-infected individuals.

One of these studies (Diepolder *et al.*, *J. Virol.* 71:6011-6019, 1997), identified peptide F98.05, which spans residues 1248-1261 of the NS3 protein, as an immunodominant CD4+ T-cell epitope that was recognized by 14/23 NS3-specific CD4+ T-cell clones from 4/5 patients with acute hepatitis C infection. This epitope, shown above to be an HLA-DR cross-reactive binder (see Table XXXIV), was capable of being presented to helper CD4+ T cells by multiple HLA molecules (DR4, DR11, DR12, DR13, and DR16). Two other peptides, Pape 22 and Pape 29, were also recognized by CD4+ T cell clones, although, in a more limited context; correspondingly, neither of these peptides are DR-cross-reactive binders.

By direct peripheral blood T cell stimulation and by fine specificity analysis of HCV-specific T-cell lines and clones, studies done in collaboration with Ferrari's group identified 6 immunodominant epitopes, including one also identified in the Pape collaboration, that are derived from conserved regions of the core, NS3, and NS4 proteins. These epitopes were also found to be cross-reactive, being presented to T cells in the context of different Class II molecules. Three of the 6 epitopes, F98.04 (F134.03), F134.05 and F134.08, are cross-reactive HLA-DR binders (see Table XXXIV).

In conclusion, the immunogenicity of 8 epitopes derived from conserved regions of the HCV genome has been demonstrated. Three of these epitopes (F98.05, F134.05, and F134.08; see Table XXXIV) are broadly cross-reactive HLA-DR binding peptides.

Example 7. Calculation of phenotypic frequencies of HLA-supertypes in various ethnic backgrounds to determine breadth of population coverage

This example illustrates the assessment of the breadth of population coverage of a vaccine composition comprised of multiple epitopes comprising multiple supermotifs and/or motifs.

In order to analyze population coverage, gene frequencies of HLA alleles were determined. Gene frequencies for each HLA allele were calculated from antigen or allele frequencies utilizing the binomial distribution formulae  $gf=1-(\text{SQRT}(1-af))$  (see, e.g., Sidney *et al.*, *Human Immunol.* 45:79-93, 1996). To obtain overall phenotypic frequencies, cumulative gene frequencies were calculated, and the cumulative antigen frequencies derived by the use of the inverse formula  $[af=1-(1-Cgf)^2]$ .

Where frequency data was not available at the level of DNA typing, correspondence to the serologically defined antigen frequencies was assumed. To obtain total potential supertype population coverage no linkage disequilibrium was assumed, and only alleles confirmed to belong to each of the supertypes were included (minimal estimates). Estimates of total potential coverage achieved by inter-loci combinations were made by adding to the A coverage the proportion of the non-A covered population that could be expected to be covered by the B alleles considered (e.g., total=A+B\*(1-A)). Confirmed members of the A3-like supertype are A3, A11, A31, A\*3301, and A\*6801. Although the A3-like supertype may also include A34, A66, and A\*7401, these alleles were not included in overall frequency calculations. Likewise, confirmed members of the A2-like supertype family are A\*0201, A\*0202, A\*0203, A\*0204, A\*0205, A\*0206, A\*0207, A\*6802, and A\*6901. Finally, the B7-like supertype-confirmed alleles are: B7, B\*3501-03, B51, B\*5301, B\*5401, B\*5501-2, B\*5601, B\*6701, and B\*7801 (potentially also B\*1401, B\*3504-06, B\*4201, and B\*5602).

Population coverage achieved by combining the A2-, A3- and B7-supertypes is approximately 86% in five major ethnic groups (see Table XXI). Coverage may be extended by including peptides bearing the A1 and A24 motifs. On average, A1 is present in 12% and A24 in 29% of the population across five different major ethnic groups (Caucasian, North American Black, Chinese, Japanese, and Hispanic). Together, these alleles are represented with an average frequency of 39% in these same ethnic populations. The total coverage across the major ethnicities when A1 and A24 are combined with the coverage of the A2-, A3- and B7-supertype alleles is >95%. An analogous approach can be used to estimate population coverage achieved with combinations of class II motif-bearing epitopes.

#### *Summary of candidate HLA class I and class II epitopes*

In summary, on the basis of the data presented in the above examples, 26 CTL candidate peptide epitopes derived from conserved regions of the HCV virus have been

identified (Table XXXVIa). These include twelve HLA-A2 supermotif-bearing epitopes, eight HLA-A3 supermotif-bearing epitopes, and one HLA-B7 supermotif-bearing epitope, each capable of binding to multiple A2-, A3-, or B7-supertype molecules, and immunogenic in HLA transgenic mice or antigenic for human PBL (with the exception of 5 peptide 29.0035/1260.04). Additional epitopes not evaluated for immunogenicity are also included. They are an additional B7-supermotif-bearing epitope and two HLA-A1 and one HLA-A24 high-affinity binding peptides. A known HLA-A31 restricted epitope (VGIYLLPNR), which also binds HLA-A33, is also set out in Table XXXVIa and is useful in combination with other Class I or Class II epitopes.

10 With these 26 CTL epitopes (as disclosed herein and from the art), average population coverage, (*i.e.*, recognition of at least one HCV epitope), is predicted to be greater than 95% in each of five major ethnic populations. The potential redundancy of coverage afforded by 25 of these epitopes (the peptide 24.0086 was not included) was estimated using the game theory Monte Carlo simulation analysis, which is known in the 15 art (see *e.g.*, Osborne, M.J. and Rubinstein, A. "A course in game theory" MIT Press, 1994). As shown in Figure 1, it is estimated that 90% of the individuals in a population comprised of the Caucasian, North American Black, Japanese, Chinese, and Hispanic ethnic groups would recognize 2 or more of the candidate epitopes described herein.

20 A list of HCV-derived HTL epitopes that would be preferred for use in the design of minigene constructs or other vaccine formulations is summarized in Table XXXVIb. As shown, 9 different peptide-binding regions have been identified which bind multiple HLA-DR molecules or bind HLA-DR3. (In the case of the NS4 1914-1935 region, the longer peptide, F134.08, recognized by patients, was chosen over the shorter peptide, 1283.44. The longer peptide essentially incorporates the shorter peptide, and also binds 25 additional DR molecules that the shorter peptide does not bind.) Three of these peptides have been recognized as dominant epitopes in HCV infected patients.

25 It is estimated that each of 10 common DR molecules recognizing the DR supermotif, and DR3, are covered by a minimum of 2 epitopes. Correspondingly, the total estimated population coverage represented by this panel of epitopes is in excess of 30 91% in each of the 5 major ethnic populations (Table XXXVII).

Example 8: Recognition Of Generation Of Endogenous Processed Antigens After Priming

This example determines that CTL induced by native or analogued peptide epitopes identified and selected as described in Examples 1-6 recognize endogenously synthesized, *i.e.*, native antigens.

Effector cells isolated from transgenic mice that are immunized with peptide epitopes as in Example 3, for example HLA-A2 supermotif-bearing epitopes, are re-stimulated *in vitro* using peptide-coated stimulator cells. Six days later, effector cells are assayed for cytotoxicity and the cell lines that contain peptide-specific cytotoxic activity 10 are further re-stimulated. An additional six days later, these cell lines are tested for cytotoxic activity on <sup>51</sup>Cr labeled Jurkat-A2.1/K<sup>b</sup> target cells in the absence or presence of peptide, and also tested on <sup>51</sup>Cr labeled target cells bearing the endogenously synthesized antigen, *i.e.* cells that are stably transfected with HCV expression vectors.

The result will demonstrate that CTL lines obtained from animals primed with 15 peptide epitope recognize endogenously synthesized HCV antigen. The choice of transgenic mouse model to be used for such an analysis depends upon the epitope(s) that is being evaluated. In addition to HLA-A\*0201/K<sup>b</sup> transgenic mice, several other transgenic mouse models including mice with human A11, which may also be used to evaluate A3 epitopes, and B7 alleles have been characterized and others (*e.g.*, transgenic 20 mice for HLA-A1 and A24) are being developed. HLA-DR1 and HLA-DR3 mouse models have also been developed, which may be used to evaluate HLA epitopes.

Example 9: Activity Of CTL-HTML Conjugated Epitopes In Transgenic Mice

This example illustrates the induction of CTLs and HTLs in transgenic mice by 25 use of an HCV CTL/HTL peptide conjugate whereby the vaccine composition comprises peptides administered to an HCV-infected patient or an individual at risk for HCV. The peptide composition can comprise multiple CTL and/or HTL epitopes. This analysis demonstrates enhanced immunogenicity that can be achieved by inclusion of one or more HTL epitopes in a vaccine composition. Such a peptide composition can comprise a 30 lipidated HTL epitope conjugated to a preferred CTL epitope containing, for example, at least one CTL epitope selected from Table XXVI-XXIX, or an analog of that epitope. The HTL epitope is, for example, selected from Table XXXII.

Lipopeptide preparation: Lipopeptides are prepared by coupling the appropriate fatty acid to the amino terminus of the resin bound peptide. A typical procedure is as

follows: A dichloromethane solution of a four-fold excess of a pre-formed symmetrical anhydride of the appropriate fatty acid is added to the resin and the mixture is allowed to react for two hours. The resin is washed with dichloromethane and dried. The resin is then treated with trifluoroacetic acid in the presence of appropriate scavengers [e.g. 5% 5 (v/v) water] for 60 minutes at 20°C. After evaporation of excess trifluoroacetic acid, the crude peptide is washed with diethyl ether, dissolved in methanol and precipitated by the addition of water. The peptide is collected by filtration and dried.

Immunization procedures: Immunization of transgenic mice is performed as described (Alexander *et al.*, *J. Immunol.* 159:4753-4761, 1997). For example, A2/K<sup>b</sup> 10 mice, which are transgenic for the human HLA A2.1 allele and are useful for the assessment of the immunogenicity of HLA-A\*0201 motif- or HLA-A2 supermotif-bearing epitopes, are primed subcutaneously (base of the tail) with 0.1 ml of peptide conjugate formulated in saline, or DMSO/saline. Seven days after priming, splenocytes obtained from these animals are restimulated with syngenic irradiated LPS-activated 15 lymphoblasts coated with peptide.

Cell lines: Target cells for peptide-specific cytotoxicity assays are Jurkat cells transfected with the HLA-A2.1/K<sup>b</sup> chimeric gene (e.g., Vitiello *et al.*, *J. Exp. Med.* 173:1007, 1991)

*In vitro* CTL activation: One week after priming, spleen cells (30x10<sup>6</sup> cells/flask) 20 are co-cultured at 37°C with syngeneic, irradiated (3000 rads), peptide coated lymphoblasts (10x10<sup>6</sup> cells/flask) in 10 ml of culture medium/T25 flask. After six days, effector cells are harvested and assayed for cytotoxic activity.

Assay for cytotoxic activity: Target cells (1.0 to 1.5x10<sup>6</sup>) are incubated at 37°C in the presence of 200 µl of <sup>51</sup>Cr. After 60 minutes, cells are washed three times and 25 resuspended in R10 medium. Peptide is added where required at a concentration of 1 µg/ml. For the assay, 10<sup>4</sup> <sup>51</sup>Cr-labeled target cells are added to different concentrations of effector cells (final volume of 200 µl) in U-bottom 96-well plates. After a 6 hour incubation period at 37°C, a 0.1 ml aliquot of supernatant is removed from each well and radioactivity is determined in a Micromedic automatic gamma counter. The percent 30 specific lysis is determined by the formula: percent specific release = 100 x (experimental release - spontaneous release)/(maximum release - spontaneous release). To facilitate comparison between separate CTL assays run under the same conditions, % <sup>51</sup>Cr release data is expressed as lytic units/10<sup>6</sup> cells. One lytic unit is arbitrarily defined as the number of effector cells required to achieve 30% lysis of 10,000 target cells in a 6

hour  $^{51}\text{Cr}$  release assay. To obtain specific lytic units/ $10^6$ , the lytic units/ $10^6$  obtained in the absence of peptide is subtracted from the lytic units/ $10^6$  obtained in the presence of peptide. For example, if 30%  $^{51}\text{Cr}$  release is obtained at the effector (E): target (T) ratio of 50:1 (i.e.,  $5 \times 10^5$  effector cells for 10,000 targets) in the absence of peptide and 5:1 (i.e., 5  $\times 10^4$  effector cells for 10,000 targets) in the presence of peptide, the specific lytic units would be:  $[(1/50,000)-(1/500,000)] \times 10^6 = 18 \text{ LU}$ .

The results are analyzed to assess the magnitude of the CTL responses of animals injected with the immunogenic CTL/HTL conjugate vaccine preparation and are compared to the magnitude of the CTL response achieved using the CTL epitope as outlined in Example 3. Analyses similar to this may be performed to evaluate the immunogenicity of peptide conjugates containing multiple CTL epitopes and/or multiple HTL epitopes. In accordance with these procedures it is found that a CTL response is induced, and concomitantly that an HTL response is induced upon administration of such compositions.

15

Example 10. Selection of CTL and HTL epitopes for inclusion in an HCV-specific vaccine.

This example illustrates the procedure for the selection of peptide epitopes for vaccine compositions of the invention. The peptides in the composition can be in the form of a nucleic acid sequence, either single or one or more sequences (i.e., minigene) that encodes peptide(s), or may be single and/or polyepitopic peptides.

Epitopes are selected which, upon administration, mimic immune responses that have been observed to be correlated with tumor clearance. For example, vaccine can include 3-4 epitopes that come from at least one HCV antigen region. Epitopes from one region can be used in combination with epitopes from one or more additional HCV antigen regions. Analogs of epitopes can also be selected for inclusion in the vaccine.

Epitopes are often selected that have a binding affinity of an  $\text{IC}_{50}$  of 500 nM or less for an HLA class I molecule, or for class II, an  $\text{IC}_{50}$  of 1000 nM or less.

Sufficient supermotif bearing peptides, or a sufficient array of allele-specific motif bearing peptides, are selected to give broad population coverage. For example, epitopes are selected to provide at least 80% population coverage. A Monte Carlo analysis, a statistical evaluation known in the art, can be employed to assess breadth, or redundancy, of population coverage.

When creating a polyepitopic compositions, *e.g.* a minigene, it is typically desirable to generate the smallest peptide possible that encompasses the epitopes of interest. The principles employed are similar, if not the same, as those employed when selecting a peptide comprising nested epitopes. Additionally, however, upon

5 determination of the nucleic acid sequence to be provided as a minigene, the peptide sequence encoded thereby is analyzed to determine whether any "junctional epitopes" have been created. A junctional epitope is a potential HLA binding epitope, as predicted, *e.g.*, by motif analysis. Junctional epitopes are generally to be avoided because the recipient may bind to an HLA molecule and generate an immune response to that epitope,

10 which is not present in a native protein sequence.

Peptide epitopes for inclusion in vaccine compositions are, for example, selected from those listed in Tables XXVI-XXIX and Table XXXII. A vaccine composition comprised of selected peptides, when administered, is safe, efficacious, and elicits an immune response similar in magnitude of an immune response that clears an acute HCV

15 infection.

Example 11: Construction of Minigene Multi-Epitope DNA Plasmids

This example provides guidance for the construction of a minigene expression plasmid. Minigene plasmids may, of course, contain various configurations of CTL and/or HTL epitopes or epitope analogs as described herein. Examples of the

20 construction and evaluation of expression plasmids are described, for example, in co-pending U.S.S.N. 09/311,784 filed 5/13/99. An example of such a plasmid for the expression of HCV epitopes is shown in Figure 2, which illustrates the orientation of HCV peptide epitopes in a minigene construct.

A minigene expression plasmid may include multiple CTL and HTL peptide epitopes. In the present example, HLA-A2, -A3, -B7 supermotif-bearing peptide epitopes and HLA-A1 and -A24 motif-bearing peptide epitopes are used in conjunction with DR supermotif-bearing epitopes and/or DR3 epitopes (Figure 2). Preferred epitopes are identified, for example, in Tables XXVI-XXIX and XXXII. HLA class I supermotif or

25 motif-bearing peptide epitopes derived from multiple HCV antigens, *e.g.*, the core, NS4, NS3, NS5, NS1/E2, are selected such that multiple supermotifs/motifs are represented to ensure broad population coverage. Similarly, HLA class II epitopes are selected from multiple HCV antigens to provide broad population coverage, *i.e.* both HLA DR-1-4-7

30 supermotif-bearing epitopes and HLA DR-3 motif-bearing epitopes are selected for

inclusion in the minigene construct. The selected CTL and HTL epitopes are then incorporated into a minigene for expression in an expression vector.

This example illustrates the methods to be used for construction of such a minigene-bearing expression plasmid. Other expression vectors that may be used for 5 minigene compositions are available and known to those of skill in the art.

The minigene DNA plasmid contains a consensus Kozak sequence and a consensus murine kappa Ig-light chain signal sequence followed by CTL and/or HTL epitopes selected in accordance with principles disclosed herein. The sequence encodes an open reading frame fused to the Myc and His antibody epitope tag coded for by the 10 pcDNA 3.1 Myc-His vector.

Overlapping oligonucleotides, for example eight oligonucleotides, averaging approximately 70 nucleotides in length with 15 nucleotide overlaps, are synthesized and HPLC-purified. The oligonucleotides encode the selected peptide epitopes as well as appropriate linker nucleotides, Kozak sequence, and signal sequence. The final 15 multiepitope minigene is assembled by extending the overlapping oligonucleotides in three sets of reactions using PCR. A Perkin/Elmer 9600 PCR machine is used and a total of 30 cycles are performed using the following conditions: 95°C for 15 sec, annealing temperature (5° below the lowest calculated Tm of each primer pair) for 30 sec, and 72°C for 1 min.

20 For the first PCR reaction, 5 µg of each of two oligonucleotides, *i.e.*, an amplification primer pair, are annealed and extended: Oligonucleotides 1+2, 3+4, 5+6, and 7+8 are combined in 100 µl reactions containing *Pfu* polymerase buffer (1x= 10 mM KCL, 10 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 20 mM Tris-chloride, pH 8.75, 2 mM MgSO<sub>4</sub>, 0.1% Triton X-100, 100 µg/ml BSA), 0.25 mM each dNTP, and 2.5 U of *Pfu* polymerase. The full-length dimer products are gel-purified, and two reactions containing the product of 1+2 and 3+4, and the product of 5+6 and 7+8 are mixed, annealed, and extended for 10 cycles. Half of the two reactions are then mixed, and 5 cycles of annealing and extension carried out before flanking primers are added to amplify the full length product for 25 additional cycles. The full-length product is gel-purified and cloned into pCR-blunt 25 (Invitrogen) and individual clones are screened by sequencing.

Example 12. The plasmid construct and the degree to which it induces immunogenicity.

The degree to which the plasmid construct prepared using the methodology outlined in Example 11 is able to induce immunogenicity is evaluated through *in vivo*

injections into mice and subsequent *in vitro* assessment of CTL and HTL activity, which are analysed using cytotoxicity and proliferation assays, respectively, as detailed *e.g.*, in U.S.S.N. 09/311,784 filed 5/13/99 and Alexander *et al.*, *Immunity* 1:751-761, 1994. For example, to assess the capacity of a pMin minigene construct that contains HLA-A2

5 supermotif epitopes to induce CTLs *in vivo*, HLA-A2.1/K<sup>b</sup> transgenic mice are immunized intramuscularly with 100 µg of naked cDNA. As a means of comparing the level of CTLs induced by cDNA immunization, a control group of animals is also immunized with an actual peptide composition that comprises multiple epitopes synthesized as a single polypeptide as they would be encoded by the minigene.

10 Splenocytes from immunized animals are stimulated twice with each of the respective compositions (peptide epitopes encoded in the minigene or the polyepitopic peptide), then assayed for peptide-specific cytotoxic activity in a <sup>51</sup>Cr release assay. The results indicate the magnitude of the CTL response directed against the A3-restricted epitope, thus indicating the *in vivo* immunogenicity of the minigene vaccine and  
15 polyepitopic vaccine. It is, therefore, found that the minigene elicits immune responses directed toward the HLA-A2 supermotif peptide epitopes as does the polyepitopic peptide vaccine. A similar analysis is also performed using other HLA-A3 and HLA-B7 transgenic mouse models to assess CTL induction by HLA-A3 and HLA-B7 motif or supermotif epitopes.

20 To assess the capacity of a class II epitope encoding minigene to induce HTLs *in vivo*, I-A<sup>b</sup> restricted mice, for example, are immunized intramuscularly with 100 µg of plasmid DNA. As a means of comparing the level of HTLs induced by DNA immunization, a group of control animals is also immunized with an actual peptide composition emulsified in complete Freund's adjuvant.

25 CD4+ T cells, *i.e.* HTLs, are purified from splenocytes of immunized animals and stimulated with each of the respective compositions (peptides encoded in the minigene). The HTL response is measured using a <sup>3</sup>H-thymidine incorporation proliferation assay, (*see, e.g.*, Alexander *et al.* *Immunity* 1:751-761, 1994). the results indicate the magnitude of the HTL response, thus demonstrating the *in vivo* immunogenicity of the minigene.

30 Alternatively, plasmid constructs can be evaluated *in vitro* by testing for epitope presentation by APC following transduction or transfection of the APC with an epitope-expressing nucleic acid construct. Such a study determines "antigenicity" and allows the use of human APC. The assay determines the ability of the epitope to be presented by the

APC in a context that is recognized by a T cell by quantifying the density of epitope-HLA class I complexes on the cell surface. Quantitation can be performed by directly measuring the amount of peptide eluted from the APC (see, e.g., Sijts *et al.*, *J. Immunol.* 156:683-692, 1996; Demotz *et al.*, *Nature* 342:682-684, 1989); or the number of peptide-HLA class I complexes can be estimated by measuring the amount of lysis or lymphokine release induced by infected or transfected target cells, and then determining the concentration of peptide necessary to obtain equivalent levels of lysis or lymphokine release (see, e.g., Kageyama *et al.*, *J. Immunol.* 154:567-576, 1995).

10 **Example 13: Peptide Composition for Prophylactic Uses**

Vaccine compositions of the present invention are used to prevent HCV infection in persons who are at risk for such infection. For example, a polyepitopic peptide epitope composition (or a nucleic acid comprising the same) containing multiple CTL and HTL epitopes such as those selected in Examples 9 and/or 10, which are also selected to target greater than 80% of the population, is administered to individuals at risk for HCV infection. The composition is provided as a single lipidated polypeptide that encompasses multiple epitopes. The vaccine is administered in an aqueous carrier comprised of Freunds Incomplete Adjuvant. The dose of peptide for the initial immunization is from about 1 to about 50,000 µg, generally 100-5,000 µg, for a 70 kg patient. The initial administration of vaccine is followed by booster dosages at 4 weeks followed by evaluation of the magnitude of the immune response in the patient, by techniques that determine the presence of epitope-specific CTL populations in a PBMC sample. Additional booster doses are administered as required. The composition is found to be both safe and efficacious as a prophylaxis against HCV infection.

25 Alternatively, the polyepitopic peptide composition can be administered as a nucleic acid in accordance with methodologies known in the art and disclosed herein.

**Example 14: Polyepitopic Vaccine Compositions Derived from Native HCV Sequences**

A native HCV polyprotein sequence is screened, preferably using computer algorithms defined for each class I and/or class II supermotif or motif, to identify "relatively short" regions of the polyprotein that comprise multiple epitopes and is preferably less in length than an entire native antigen. This relatively short sequence that contains multiple distinct, even overlapping, epitopes is selected and used to generate a minigene construct. The construct is engineered to express the peptide, which

corresponds to the native protein sequence. The "relatively short" peptide is generally less than 250 amino acids in length, often less than 100 amino acids in length, preferably less than 75 amino acids in length, and more preferably less than 50 amino acids in length. The protein sequence of the vaccine composition is selected because it has

5 maximal number of epitopes contained within the sequence, *i.e.*, it has a high concentration of epitopes. As noted herein, epitope motifs may be nested or overlapping (*i.e.*, frame shifted relative to one another). For example, with frame shifted overlapping epitopes, two 9-mer epitopes and one 10-mer epitope can be present in a 10 amino acid peptide. Such a vaccine composition is administered for therapeutic or prophylactic

10 purposes.

The vaccine composition will preferably include, for example, three CTL epitopes and at least one HTL epitope from an HCV antigen. This polyepitopic native sequence is administered either as a peptide or as a nucleic acid sequence which encodes the peptide. Alternatively, an analog can be made of this native sequence, whereby one or more of the

15 epitopes comprise substitutions that alter the cross-reactivity and/or binding affinity properties of the polyepitopic peptide.

The embodiment of this example provides for the possibility that an as yet undiscovered aspect of immune system processing will apply to the native nested sequence and thereby facilitate the production of therapeutic or prophylactic immune

20 response-inducing vaccine compositions. Additionally such an embodiment provides for the possibility of motif-bearing epitopes for an HLA makeup that is presently unknown. Furthermore, this embodiment (absent analogs) directs the immune response to multiple peptide sequences that are actually present in native HCV antigens thus avoiding the need to evaluate any junctional epitopes. Lastly, the embodiment provides an economy of

25 scale when producing nucleic acid vaccine compositions.

Related to this embodiment, computer programs can be derived in accordance with principles in the art, which identify in a target sequence, the greatest number of epitopes per sequence length.

30 Example 15. Polyepitopic Vaccine Compositions Directed To Multiple Diseases

The HCV peptide epitopes of the present invention are used in conjunction with peptide epitopes from target antigens related to one or more other diseases, to create a vaccine composition that is useful for the prevention or treatment of HCV as well as the

one or more other disease(s). Examples of the other diseases include, but are not limited to, HIV, and HBV.

For example, a polyepitopic peptide composition comprising multiple CTL and HTL epitopes that target greater than 98% of the population may be created for

5 administration to individuals at risk for both HCV and HIV infection. The composition can be provided as a single polypeptide that incorporates the multiple epitopes from the various disease-associated sources, or can be administered as a composition comprising one or more discrete epitopes.

10 Example 16. Use of peptides to evaluate an immune response

Peptides of the invention may be used to analyze an immune response for the presence of specific CTL or HTL populations directed to a prostate cancer-associated antigen. Such an analysis may be performed using multimeric complexes as described, e.g., by Ogg *et al.*, *Science* 279:2103-2106, 1998 and Greten *et al.*, *Proc. Natl. Acad. Sci. USA* 95:7568-7573, 1998. In the following example, peptides in accordance with the invention are used as a reagent for diagnostic or prognostic purposes, not as an immunogen.

In this example, highly sensitive human leukocyte antigen tetrameric complexes ("tetramers") are used for a cross-sectional analysis of, for example, HCV HLA-A\*0201-specific CTL frequencies from HLA A\*0201-positive individuals at different stages of disease or following immunization using an HCV peptide containing an A\*0201 motif. Tetrameric complexes are synthesized as described (Musey *et al.*, *N. Engl. J. Med.* 337:1267, 1997). Briefly, purified HLA heavy chain (A\*0201 in this example) and  $\beta$ 2-microglobulin are synthesized by means of a prokaryotic expression system. The heavy chain is modified by deletion of the transmembrane-cytosolic tail and COOH-terminal addition of a sequence containing a BirA enzymatic biotinylation site. The heavy chain,  $\beta$ 2-microglobulin, and peptide are refolded by dilution. The 45-kD refolded product is isolated by fast protein liquid chromatography and then biotinylated by BirA in the presence of biotin (Sigma, St. Louis, Missouri), adenosine 5' triphosphate and magnesium. Streptavidin-phycoerythrin conjugate is added in a 1:4 molar ratio, and the tetrameric product is concentrated to 1 mg/ml. The resulting product is referred to as tetramer-phycoerythrin.

For the analysis of patient blood samples, approximately one million PBMCs are centrifuged at 300g for 5 minutes and resuspended in 50  $\mu$ l of cold phosphate-buffered saline. Tri-color analysis is performed with the tetramer-phycerythrin, along with anti-CD8-Tricolor, and anti-CD38. The PBMCs are incubated 5 with tetramer and antibodies on ice for 30 to 60 min and then washed twice before formaldehyde fixation. Gates are applied to contain >99.98% of control samples. Controls for the tetramers include both A\*0201-negative individuals and A\*0201-positive uninfected donors. The percentage of cells stained with the tetramer is then determined by flow cytometry. The results indicate the number of cells in the PBMC sample that 10 contain epitope-restricted CTLs, thereby readily indicating the extent of immune response to the HCV epitope, and thus the stage of HCV infection or exposure to a vaccine that elicits a protective or therapeutic response.

Example 17: Use of Peptide Epitopes to Evaluate Recall Responses

15 The peptide epitopes of the invention are used as reagents to evaluate T cell responses, such as acute or recall responses, in patients. Such an analysis may be performed on patients who have recovered from infection, who are chronically infected with HCV, or who have been vaccinated with an HCV vaccine.

20 For example, the class I restricted CTL response of persons who have been vaccinated may be analyzed. The vaccine may be any HCV vaccine. PBMC are collected from vaccinated individuals and HLA typed. Appropriate peptide epitopes of the invention that are preferably highly conserved and, optimally, bear supermotifs to provide cross-reactivity with multiple HLA supertype family members, are then used for analysis of samples derived from individuals who bear that HLA type.

25 PBMC from vaccinated individuals are separated on Ficoll-Histopaque density gradients (Sigma Chemical Co., St. Louis, MO), washed three times in HBSS (GIBCO Laboratories), resuspended in RPMI-1640 (GIBCO Laboratories) supplemented with L-glutamine (2mM), penicillin (50U/ml), streptomycin (50  $\mu$ g/ml), and Hepes (10mM) containing 10% heat-inactivated human AB serum (complete RPMI) and plated using 30 microculture formats. A synthetic peptide comprising an epitope of the invention is added at 10  $\mu$ g/ml to each well and HBV core 128-140 epitope is added at 1  $\mu$ g/ml to each well as a source of T cell help during the first week of stimulation.

In the microculture format,  $4 \times 10^5$  PBMC are stimulated with peptide in 8 replicate cultures in 96-well round bottom plate in 100  $\mu$ l/well of complete RPMI. On

days 3 and 10, 100 ml of complete RPMI and 20 U/ml final concentration of rIL-2 are added to each well. On day 7 the cultures are transferred into a 96-well flat-bottom plate and restimulated with peptide, rIL-2 and 10<sup>5</sup> irradiated (3,000 rad) autologous feeder cells. The cultures are tested for cytotoxic activity on day 14. A positive CTL response 5 requires two or more of the eight replicate cultures to display greater than 10% specific <sup>51</sup>Cr release, based on comparison with uninfected control subjects as previously described (Rehermann, *et al.*, *Nature Med.* 2:1104,1108, 1996; Rehermann *et al.*, *J. Clin. Invest.* 97:1655-1665, 1996; and Rehermann *et al.* *J. Clin. Invest.* 98:1432-1440, 1996).

Target cell lines are autologous and allogeneic EBV-transformed B-LCL that are 10 either purchased from the American Society for Histocompatibility and Immunogenetics (ASHI, Boston, MA) or established from the pool of patients as described (Guilhot, *et al.* *J. Virol.* 66:2670-2678, 1992).

Cytotoxicity assays are performed in the following manner. Target cells consist 15 of either allogeneic HLA-matched or autologous EBV-transformed B lymphoblastoid cell line that are incubated overnight with the synthetic peptide epitope of the invention at 10  $\mu$ M, and labeled with 100  $\mu$ Ci of <sup>51</sup>Cr (Amersham Corp., Arlington Heights, IL) for 1 hour after which they are washed four times with HBSS.

Cytolytic activity is determined in a standard 4-h, split well <sup>51</sup>Cr release assay 20 using U-bottomed 96 well plates containing 3,000 targets/well. Stimulated PBMC are tested at effector/target (E/T) ratios of 20-50:1 on day 14. Percent cytotoxicity is determined from the formula: 100 x [(experimental release-spontaneous release)/maximum release-spontaneous release)]. Maximum release is determined by lysis of targets by detergent (2% Triton X-100; Sigma Chemical Co., St. Louis, MO). Spontaneous release is <25% of maximum release for all experiments.

25 The results of such an analysis indicate the extent to which HLA-restricted CTL populations have been stimulated by previous exposure to HCV or an HCV vaccine.

The class II restricted HTL responses may also be analyzed. Purified PBMC are cultured in a 96-well flat bottom plate at a density of 1.5x10<sup>5</sup> cells/well and are stimulated with 10  $\mu$ g/ml synthetic peptide, whole antigen, or PHA. Cells are routinely plated in 30 replicates of 4-6 wells for each condition. After seven days of culture, the medium is removed and replaced with fresh medium containing 10U/ml IL-2. Two days later, 1  $\mu$ Ci <sup>3</sup>H-thymidine is added to each well and incubation is continued for an additional 18 hours. Cellular DNA is then harvested on glass fiber mats and analyzed for <sup>3</sup>H-thymidine

incorporation. Antigen-specific T cell proliferation is calculated as the ratio of  $^3\text{H}$ -thymidine incorporation in the presence of antigen divided by the  $^3\text{H}$ -thymidine incorporation in the absence of antigen.

5 Example 18: Induction Of Specific CTL Response In Humans

A human clinical trial for an immunogenic composition comprising CTL and HTL epitopes of the invention is set up as an IND Phase I, dose escalation study and carried out as a randomized, double-blind, placebo-controlled trial. Such a trial is designed, for example, as follows:

10 A total of about 27 subjects are enrolled and divided into 3 groups:

Group I: 3 subjects are injected with placebo and 6 subjects are injected with 5  $\mu\text{g}$  of peptide composition;

Group II: 3 subjects are injected with placebo and 6 subjects are injected with 50  $\mu\text{g}$  peptide composition;

15 Group III: 3 subjects are injected with placebo and 6 subjects are injected with 500  $\mu\text{g}$  of peptide composition.

After 4 weeks following the first injection, all subjects receive a booster inoculation at the same dosage.

20 The endpoints measured in this study relate to the safety and tolerability of the peptide composition as well as its immunogenicity. Cellular immune responses to the peptide composition are an index of the intrinsic activity of this the peptide composition, and can therefore be viewed as a measure of biological efficacy. The following summarize the clinical and laboratory data that relate to safety and efficacy endpoints.

25 Safety: The incidence of adverse events is monitored in the placebo and drug treatment group and assessed in terms of degree and reversibility.

Evaluation of Vaccine Efficacy: For evaluation of vaccine efficacy, subjects are bled before and after injection. Peripheral blood mononuclear cells are isolated from fresh heparinized blood by Ficoll-Hypaque density gradient centrifugation, aliquoted in freezing media and stored frozen. Samples are assayed for CTL and HTL activity.

30 The vaccine is found to be both safe and efficacious.

Example 19: Phase II Trials In Patients Infected With HCV

Phase II trials are performed to study the effect of administering the CTL-HTL peptide compositions to patients having chronic HCV infection. The main objectives of

the trials are to determine an effective dose and regimen for inducing CTLs in chronically infected HCV patients, to establish the safety of inducing a CTL and HTL response in these patients, and to see to what extent activation of CTLs improves the clinical picture of chronically infected CTL patients, as manifested by a transient flare in alanine 5 aminotransferase (ALT), normalization of ALT, and reduction in HCV DNA. Such a study is designed, for example, as follows:

The studies are performed in multiple centers. The trial design is an open-label, uncontrolled, dose escalation protocol wherein the peptide composition is administered as a single dose followed six weeks later by a single booster shot of the same dose. The 10 dosages are 50, 500 and 5,000 micrograms per injection. Drug-associated adverse effects (severity and reversibility) are recorded.

There are three patient groupings. The first group is injected with 50 micrograms of the peptide composition and the second and third groups with 500 and 5,000 15 micrograms of peptide composition, respectively. The patients within each group range in age from 21-65, include both males and females, and represent diverse ethnic backgrounds. All of them are infected with HCV for over five years and are HIV, HBV and delta hepatitis virus (HDV) negative, but have positive levels of HCV antigen.

The magnitude and incidence of ALT flares and the levels of HCV DNA in the blood are monitored to assess the effects of administering the peptide compositions. The 20 levels of HCV DNA in the blood are an indirect indication of the progress of treatment. The vaccine composition is found to be both safe and efficacious in the treatment of chronic HCV infection.

Example 20. Induction of CTL Responses Using a Prime Boost Protocol

25 A prime boost protocol can also be used for the administration of the vaccine to humans. Such a vaccine regimen may include an initial administration of, for example, naked DNA followed by a boost using recombinant virus encoding the vaccine, or recombinant protein/polypeptide or a peptide mixture administered in an adjuvant.

30 For example, the initial immunization may be performed using an expression vector, such as that constructed in Example 11, in the form of naked nucleic acid administered IM (or SC or ID) in the amounts of 0.5-5 mg at multiple sites. The nucleic acid (0.1 to 1000  $\mu$ g) can also be administered using a gene gun. Following an incubation period of 3-4 weeks, a booster dose is administered. The booster can, e.g., be recombinant fowlpox virus administered at a dose of  $5 \cdot 10^7$  to  $5 \times 10^9$  pfu. An alternative

recombinant virus, such as an MVA, canarypox, adenovirus, or adeno-associated virus, can also be used for the booster, or the polyepitopic protein or a mixture of the peptides can be administered. For evaluation of vaccine efficacy, patient blood samples will be obtained before immunization as well as at intervals following administration of the 5 initial vaccine and booster doses of the vaccine. Peripheral blood mononuclear cells are isolated from fresh heparinized blood by Ficoll-Hypaque density gradient centrifugation, aliquoted in freezing media and stored frozen. Samples are assayed for CTL and HTL activity.

10 Analysis of the results will indicate that a magnitude of response sufficient to achieve protective immunity or to treat HCV infection infection is generated.

Example 21. Administration of Vaccine Compositions Using Dendritic Cells

Vaccines comprising peptide epitopes of the invention may be administered using dendritic cells. In this example, the peptide-pulsed dendritic cells can be administered to 15 a patient to stimulate a CTL response *in vivo*. In this method dendritic cells are isolated, expanded, and pulsed with a vaccine comprising peptide CTL and HTL epitopes of the invention. The dendritic cells are infused back into the patient to elicit CTL and HTL responses *in vivo*. The induced CTL and HTL then destroy (CTL) or facilitate destruction 20 (HTL) of the specific target HCV-infected cells that bear the proteins from which the epitopes in the vaccine are derived.

Alternatively, *Ex vivo* CTL or HTL responses to a particular tumor-associated antigen can be induced by incubating in tissue culture the patient's, or genetically compatible, CTL or HTL precursor cells together with a source of antigen-presenting cells, such as dendritic cells, and the appropriate immunogenic peptides. After an 25 appropriate incubation time (typically about 7-28 days), in which the precursor cells are activated and expanded into effector cells, the cells are infused back into the patient, where they will destroy (CTL) or facilitate destruction (HTL) of their specific target cells, *i.e.*, tumor cells.

30 Example 22: Alternative Method of Identifying Motif-Bearing Peptides

Another way of identifying motif-bearing peptides is to elute them from cells bearing defined MHC molecules. For example, EBV transformed B cell lines used for tissue typing, have been extensively characterized to determine which HLA molecules they express. In certain cases these cells express only a single type of HLA molecule.

These cells can then be infected with a pathogenic organism, *e.g.*, HCV, or transfected with nucleic acids that express the antigen of interest. Thereafter, peptides produced by endogenous antigen processing of peptides produced consequent to infection (or as a result of transfection) will bind be displayed on the cell surface. The peptides are then 5 eluted from the HLA molecules by exposure to mild acid conditions and their amino acid sequence determined, *e.g.*, by mass spectral analysis (*e.g.*, Kubo *et al.*, *J. Immunol.* 152:3913, 1994). Because, as disclosed herein, the majority of peptides that bind a particular HLA molecule are motif-bearing, this is an alternative modality for obtaining the motif-bearing peptides correlated with the particular HLA molecule expressed on the 10 cell.

Alternatively, cell lines that do not express any endogenous HLA molecules can be transfected with an expression construct encoding a single HLA allele. These cells may then be used as described, *i.e.*, they may be infected with a pathogenic organism or transfected with nucleic acid encoding an antigen of interest to isolate peptides 15 corresponding to the pathogen or antigen of interest that have been presented on the cell surface. Peptides obtained from such an analysis will bear motif(s) that correspond to binding to the single HLA allele that is expressed in the cell.

As appreciated by one in the art, one can perform a similar analysis on a cell bearing more than one HLA allele and subsequently determine peptides specific for each 20 HLA allele expressed. Moreover, one of skill would also recognize that means other than infection or transfection, such as loading with a protein antigen, can be used to provide a source of antigen to the cell.

The above examples are provided to illustrate the invention but not to limit its scope. For example, the human terminology for the Major Histocompatibility Complex, 25 namely HLA, is used throughout this document. It is to be appreciated that these principles can be extended to other species as well. Thus, other variants of the invention will be readily apparent to one of ordinary skill in the art and are encompassed by the appended claims. All publications, patents, and patent application cited herein are hereby incorporated by reference for all purposes.

TABLE I

SUPERMOTIFS	POSITION	POSITION	POSITION
	2 (Primary Anchor)	3 (Primary Anchor)	C Terminus (Primary Anchor)
A1	<b>T, I, L, V, M, S</b>		<b>F, W, Y</b>
A2	<b>L, I, V, M, A, T, Q</b>		<b>I, V, M, A, T, L</b>
A3	<b>V, S, M, A, T, L, I</b>		<b>R, K</b>
A24	<b>Y, F, W, I, V, L, M, T</b>		<b>F, I, Y, W, L, M</b>
B7	<b>P</b>		<b>V, I, L, F, M, W, Y, A</b>
B27	<b>R, H, K</b>		<b>F, Y, L, W, M, I, V, A</b>
B44	<b>E, D</b>		<b>F, W, L, I, M, V, A</b>
B58	<b>A, T, S</b>		<b>F, W, Y, L, I, V, M, A</b>
B62	<b>Q, L, I, V, M, P</b>		<b>F, W, Y, M, I, V, L, A</b>
MOTIFS			
A1	<b>T, S, M</b>		<b>Y</b>
A1		<b>D, E, A, S</b>	<b>Y</b>
A2.1	<b>L, M, V, Q, I, A, T</b>		<b>V, L, I, M, A, T</b>
A3	<b>L, M, V, I, S, A, T, F, C, G, D</b>		<b>K, Y, R, H, F, A</b>
A11	<b>V, T, M, L, I, S, A, G, N, C, D, F</b>		<b>K, R, Y, H</b>
A24	<b>Y, F, W, M</b>		<b>F, L, I, W</b>
A*3101	<b>M, V, T, A, L, I, S</b>		<b>R, K</b>
A*3301	<b>M, V, A, L, F, I, S, T</b>		<b>R, K</b>
A*6801	<b>A, V, T, M, S, L, I</b>		<b>R, K</b>
B*0702	<b>P</b>		<b>L, M, F, W, Y, A, I, V</b>
B*3501	<b>P</b>		<b>L, M, F, W, Y, I, V, A</b>
B51	<b>P</b>		<b>L, I, V, F, W, Y, A, M</b>
B*5301	<b>P</b>		<b>I, M, F, W, Y, A, L, V</b>
B*5401	<b>P</b>		<b>A, T, I, V, L, M, F, W, Y</b>

Bolded residues are preferred, italicized residues are less preferred: A peptide is considered motif-bearing if it has primary anchors at each primary anchor position for a motif or supermotif as specified in the above table.

TABLE II

## SUPERMOTIFS

		POSITION								
		1	2	3	4	5	6	7	8	C-terminus
<b>SUPERMOTIFS</b>										
A1										1° Anchor F,W,Y
										1° Anchor L,I,V,M,A,T
A2										1° Anchor L,I,V,M,A,T
A3	preferred					1° Anchor V,S,M,A,T, L,I	Y,F,W (4/5)	Y,F,W (4/5)	P (4/5)	1° Anchor R,K
	deleterious					D,E (3/5); P (5/5)	D,E (4/5)			
A24							1° Anchor Y,F,W,I,V, L,M,T			1° Anchor F,I,Y,W,I,M
B7	preferred					F,W,Y (5/5) L,I,V,M (3/5)	1° Anchor P	F,W,Y (4/5)		1° Anchor V,I,L,F,M,W,Y,A
	deleterious					D,E (3/5); P (5/5); G(4/5); A(3/5); Q,N (3/5)	D,E (3/5)	G (4/5)	Q,N (4/5)	D,E (4/5)
B27							1° Anchor R,H,K			1° Anchor F,Y,L,W,M,V,A
B44							1° Anchor E,D			1° Anchor F,W,Y,L,I,M,V,A
B58							1° Anchor A,T,S			1° Anchor F,W,Y,L,I,V,M,A
B62							1° Anchor Q,I,J,V,M, P			1° Anchor F,W,Y,M,I,V,L,A

		POSITION									
		1	2	3	4	5	6	7	8	C-terminus	
MOTIFS											
A1 9-mer	preferred	G,F,Y,W		<sup>1°</sup> Anchor S,T,M	D,E,A	Y,F,W		P	D,E,Q,N	Y,F,W	<sup>1°</sup> Anchor Y
	deleterious	D,E		R,H,K,L,I,V M,P	A	G	A				
A1 9-mer	preferred	G,R,H,K		A,S,T,C,L,I V,M, V,M,	<sup>1°</sup> Anchor D,E,A,S	G,S,T,C		A,S,T,C	L,I,V,M	D,E	<sup>1°</sup> Anchor Y
	deleterious	A		R,H,K,D,E, P,Y,F,W	D,E	P,Q,N	R,H,K	P,G	G,P		

		POSITION								
		1	2	3	4	5	6	7	8	9
A1 10-mer	preferred	Y,F,W	<sup>1°Anchor</sup> S,T,M	D,E,A,Q,N	A	Y,F,W,Q,N		P,A,S,T,C	G,D,E	P
	deleterious	G,P		R,H,K,G,L,I V,M	D,E	R,H,K	Q,N,A	R,H,K,Y,F, W	R,H,K	A
A1 10-mer	preferred	Y,F,W	S,T,C,L,I,V M	<sup>1°Anchor</sup> D,E,A,S	A	Y,F,W		P,G	G	Y,F,W
	deleterious	R,H,K		R,H,K,D,E, P,Y,F,W		P	G	P,R,H,K	Q,N	<sup>1°Anchor</sup> Y
A2.1 9-mer	preferred	Y,F,W	<sup>1°Anchor</sup> L,M,I,V,Q, A,T	Y,F,W	S,T,C	Y,F,W		A	P	<sup>1°Anchor</sup> V,L,I,M,A,T
	deleterious	D,E,P		D,E,R,K,H		R,K,H	D,E,R,K,H			
A2.1 10-mer	preferred	A,Y,F,W	<sup>1°Anchor</sup> L,M,I,V,Q, A,T	L,V,I,M	G		G			<sup>1°Anchor</sup> V,L,I,M
	deleterious	D,E,P		D,E	R,K,H,A	P	R,K,H	D,E,R, K,H	R,K,H	<sup>1°Anchor</sup> V,L,I,M,A,T

		POSITION									
		1	2	3	4	5	6	7	8	9	C-terminus
A3 preferred		R,H,K	<sup>1°</sup> Anchor L,M,V,I,S, A,T,F,C,G D	Y,F,W	P,R,H,K,Y, F,W	A	Y,F,W		P	C-terminus <sup>1°</sup> Anchor K,Y,R,H,F,A	
deleterious		D,E,P		D,E							
A11 preferred		A	<sup>1°</sup> Anchor V,T,L,M,I, S,A,G,N,C, D,F	Y,F,W	Y,F,W	A	Y,F,W	Y,F,W	P	<sup>1°</sup> Anchor K,R,Y,H	
deleterious		D,E,P					A	G			
A24 preferred 9-mer		Y,F,W,R,H,K	<sup>1°</sup> Anchor Y,F,W,M		S,T,C		Y,F,W	Y,F,W		<sup>1°</sup> Anchor F,L,I,W	
deleterious		D,E,G		D,E	G	Q,N,P	D,E,R,H,K	G	A,Q,N		
A24 preferred 10-mer			<sup>1°</sup> Anchor Y,F,W,M	P	Y,F,W,P		P			<sup>1°</sup> Anchor F,L,I,W	
deleterious			G,D,E	Q,N	R,H,K	D,E	A	Q,N	D,E,A		
A3101 preferred		R,H,K	<sup>1°</sup> Anchor M,V,T,A,L, I,S	Y,F,W	P	Y,F,W	Y,F,W	A,P	<sup>1°</sup> Anchor R,K		
deleterious		D,E,P		D,E		D,E	D,E	D,E			

		POSITION								
		1	2	3	4	5	6	7	8	9
A3301 preferred		<u>1°Anchor</u> M,V,A,L,F, I,S,T	Y,F,W							C-terminus or C-terminus 1°Anchor R,K
deleterious		G,P			D,E					
A6801 preferred		Y,F,W,S,T,C	<u>1°Anchor</u> A,V,T,M,S, L,I			Y,F,W,L,I, V,M		Y,F,W	P	1°Anchor R,K
deleterious		G,P		D,E,G		R,H,K				A
B0702 preferred		R,H,K,F,W,Y	<u>1°Anchor</u> P	R,H,K		R,H,K		R,H,K	P,A	1°Anchor L,M,F,W,Y,A, I,V
deleterious		D,E,Q,N,P		D,E,P	D,E	D,E	G,D,E	Q,N	D,E	
B3501 preferred		F,W,Y,L,I,V,M	<u>1°Anchor</u> P	F,W,Y				F,W,Y		1°Anchor L,M,F,W,Y,I, V,A
deleterious		A,G,P				G	G			

		POSITION								
		1	2	3	4	5	6	7	8	9
B51	preferred	L,I,V,M,F,W,Y	<u>P</u> <u>1°Anchor</u>	F,W,Y	S,T,C	F,W,Y	G	F,W,Y	F,W,Y	C-terminus or C-terminus 1°Anchor L,I,V,F,W, Y,A,M
	deleterious	A,G,P,D,E,R,H,K, S,T,C			D,E	G	D,E,Q,N	G,D,E		
B5301	preferred	L,I,V,M,F,W,Y	<u>P</u> <u>1°Anchor</u>	F,W,Y	S,T,C	F,W,Y	L,I,V,M,F, W,Y	F,W,Y	F,W,Y	1°Anchor I,M,F,W,Y, A,L,V
	deleterious	A,G,P,Q,N				G	R,H,K,Q,N	D,E		
B5401	preferred	F,W,Y	<u>P</u> <u>1°Anchor</u>	F,W,Y,I,V M		L,I,V,M	A,L,I,V,M	F,W,Y,A,P	F,W,Y,A,P	1°Anchor A,T,I,V,L, M,F,W,Y
	deleterious	G,P,Q,N,D,E		G,D,E,S,T,C	R,H,K,D,E	D,E	Q,N,D,G,E	D,E		

Italicized residues indicate less preferred or "tolerated" residues.  
The information in Table II is specific for 9-mers unless otherwise specified.

Table III

MOTIFS		POSITION			
1° anchor 1		2	3	4	5
DR4 preferred	F, M, Y, L, I, <i>V, W</i>	M	T	I	V, S, T, C, P, A, <i>L, I, M</i>
deleterious			W,		M, H, R,
DR1 preferred	M, F, L, I, V, <i>W, Y</i>	C	C, H	P, A, M, Q F, D	V, M, A, T, S, P, <i>L, I, C</i>
deleterious			W	A	G, D, E, G, D, E, D
DR7 preferred	M, F, L, I, V, <i>W, Y</i>	M			I, V, M, S, A, C, <i>T, P, L</i>
deleterious		C,		G,	G, R, D
DR Supermotif	M, F, L, I, V, <i>W, Y</i>				V, M, S, T, A, C, <i>P, L, I</i>
DR3 MOTIFS		1° anchor 1	2	3	4
motif a preferred	L, I, V, M, F, <i>Y</i>				D
motif b preferred	L, I, V, M, F, <i>A, Y</i>				D, N, Q, E, <i>S, T</i>
					K, R, H

Italicized residues indicate less preferred or "tolerated" residues.

**Table IV: HLA Class I Standard Peptide Binding Affinity.**

ALLEL	STANDARD PEPTIDE	SEQUENCE (SEQ ID NO:)	STANDARD BINDING AFFINITY (nM)
A*0101	944.02	YLEPAIAKY	25
A*0201	941.01	FLPSDYFPSV	5.0
A*0202	941.01	FLPSDYFPSV	4.3
A*0203	941.01	FLPSDYFPSV	10
A*0205	941.01	FLPSDYFPSV	4.3
A*0206	941.01	FLPSDYFPSV	3.7
A*0207	941.01	FLPSDYFPSV	23
A*6802	1072.34	YVIKVSARV	8.0
A*0301	941.12	KVFPYALINK	11
A*1101	940.06	AVDLYHFLK	6.0
A*3101	941.12	KVFPYALINK	18
A*3301	1083.02	STLPETYVVRR	29
A*6801	941.12	KVFPYALINK	8.0
A*2402	979.02	AYIDNYNKF	12
B*0702	1075.23	APRTLVYLL	5.5
B*3501	1021.05	FPFKYAAAF	7.2
B51	1021.05	FPFKYAAAF	5.5
B*5301	1021.05	FPFKYAAAF	9.3
B*5401	1021.05	FPFKYAAAF	10

Table V. HLA Class II Standard Peptide Binding Affinity.

Allele	Nomenclature	Standard Peptide	Sequence (SEQ ID NO:)	Binding Affinity (nM)
DRB1*0101	DR1	515.01	PKYVKQNTLKLAT	5.0
DRB1*0301	DR3	829.02	YKTIAFDEEARR	300
DRB1*0401	DR4w4	515.01	PKYVKQNTLKLAT	45
DRB1*0404	DR4w14	717.01	YARFQSQTTLKQKT	50
DRB1*0405	DR4w15	717.01	YARFQSQTTLKQKT	38
DRB1*0701	DR7	553.01	QYIKANSKFIGITE	25
DRB1*0802	DR8w2	553.01	QYIKANSKFIGITE	49
DRB1*0803	DR8w3	553.01	QYIKANSKFIGITE	1600
DRB1*0901	DR9	553.01	QYIKANSKFIGITE	75
DRB1*1101	DR5w11	553.01	QYIKANSKFIGITE	20
DRB1*1201	DR5w12	1200.05	EALIIHQLKINPYVLS	298
DRB1*1302	DR6w19	650.22	QYIKANAKFIGITE	3.5
DRB1*1501	DR2w2 $\beta$ 1	507.02	GRTQDENPVVHFFKNIV TPRTPPP	9.1
DRB3*0101	DR52a	511	NGQIGNDPNRDIL	470
DRB4*0101	DRw53	717.01	YARFQSQTTLKQKT	58
DRB5*0101	DR2w2 $\beta$ 2	553.01	QYIKANSKFIGITE	20

Table VI

HLA-supertype	Verified <sup>a</sup>	Allele-specific HLA-supertype members	Predicted <sup>b</sup>
A1	A*0101, A*2501, A*2601, A*2602, A*3201		A*0102, A*2604, A*3601, A*4301, A*8001
A2	A*0201, A*0202, A*0203, A*0204, A*0205, A*0206, A*0207, A*0209, A*0214, A*6802, A*6901		A*0208, A*0210, A*0211, A*0212, A*0213
A3	A*0301, A*1101, A*3101, A*3301, A*6801		A*0302, A*1102, A*2603, A*3302, A*3303, A*3401, A*3402, A*6601, A*6602, A*7401
A24	A*2301, A*2402, A*3001		A*2403, A*2404, A*3002, A*3003
B7	B*0702, B*0703, B*0704, B*0705, B*1508, B*3501, B*3502, B*3503, B*3503, B*3504, B*3505, B*3506, B*3507, B*3508, B*5101, B*5102, B*5103, B*5104, B*5105, B*5301, B*5401, B*5501, B*5502, B*5601, B*5602, B*6701, B*7801		B*1511, B*4201, B*5901
B27	B*1401, B*1402, B*1509, B*2702, B*2703, B*2704, B*2705, B*2706, B*3801, B*3901, B*3902, B*3701		B*2701, B*2707, B*2708, B*3802, B*3903, B*3904, B*3905, B*4801, B*4802, B*1510, B*1518, B*1503
B44	B*1801, B*1802, B*3701, B*4402, B*4403, B*4404, B*4001, B*4002, B*4006		B*4101, B*4501, B*4701, B*4901, B*5001
B58	B*5701, B*5702, B*5801, B*5802, B*1516, B*1517		
B62	B*1501, B*1502, B*1513, B*5201		B*1301, B*1302, B*1504, B*1505, B*1506, B*1507, B*1515, B*1520, B*1521, B*1512, B*1514, B*1510

a. Verified alleles include alleles whose specificity has been determined by pool sequencing analysis, peptide binding assays, or by analysis of the sequences of CTL epitopes.

b. Predicted alleles are alleles whose specificity is predicted on the basis of B and F pocket structure to overlap with the supertype specificity.

Table VII HCV A01 Super Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*0101
ATGNLPCSF	165	10	13	93	
ATLGFTAY	1265	8	14	100	
AVQWMMRLIAF	1917	11	14	100	
CTCGSSDLY	1128	9	11	79	0.3700
CTRGVAKAYDF	1190	11	11	79	
CTWMNSTGF	555	9	11	79	
CYQVDF	1462	8	12	86	
DLEVNTSW	1857	9	12	86	
ETTMRSIPVF	1207	9	12	86	
FSDYTRCF	2870	8	11	79	
FTEAMTRY	2792	8	14	100	
FTGLTHIDAHF	1567	11	13	93	
GLVYCCDHLF	1552	11	12	86	
GLSAFSLSHSY	2821	10	11	79	0.0029
GLTHIDAHF	1569	9	13	93	
GSSYGEFY	2641	8	11	79	
GTFPNAY	2063	8	11	79	
GVAGALVAF	1863	9	12	86	
GVAKAVIDF	1183	8	11	79	
GVLAALDAY	1670	9	12	86	
GVRVCEKMLY	2619	11	14	100	
GVRMEOGMY	154	11	12	86	
HHCQNVQVQY	696	11	11	79	
HMWNFGIGQY	1769	11	13	93	
HMGEGEGAYW	1910	11	11	79	
IMARKEVF	2591	8	12	86	
ITYSTYGF	1296	9	12	86	
IVDQDLY	701	8	12	86	
KSTKVNPAAY	1241	9	12	86	0.0130
KVDTLTCGF	121	10	12	86	
LIEANLW	2235	8	12	86	
LINTNGSN	414	8	11	79	
LLAPITAY	1030	8	14	100	
LLFNLGAW	1812	9	12	86	
LLSPFGSIPSW	97	11	11	79	
LSAFSLHSY	2922	9	11	79	0.8100
LSPFGSIPSW	98	10	11	79	
LTCGFADIMGY	126	11	12	86	
LTHIDAHF	1570	8	13	93	
LYDILAGY	1853	8	11	79	
MILMTIFF	2878	8	12	86	
NIVDQMY	700	9	12	86	
NUPGCSFSIF	168	10	13	93	
NTCTGIVDF	1460	10	12	86	
NTNFRPODF	14	11	11	78	

## HCV A01 Super Motif with Binding Information

Sequence	Position		No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'0101
NVQDGW	1108		9	11	79	
PIYSTYAKF	1295		10	11	79	
PNFGSYDTRCF	2667		11	11	79	
PSVAATLGF	1281		9	14	100	
PTLHGPTPLY	1621		11	11	79	
PVQDQALEF	1554		9	12	86	
PVQDQALEFW	1554		10	12	86	
QTVQFSLDPF	1465		11	12	86	
RUJGLSAF	2918		8	12	86	
RLLAPHTAY	1029		9	12	86	
RMAWDMMMMW	317		10	12	86	
RMLMLTHF	2875		9	12	86	
RMLMLTHF	2875		9	12	86	
RVCERKMY	2621		9	14	100	
RMLEGVNV	156		9	12	86	
STKVPAY	1242		8	12	86	
SVAAATLGF	1262		8	14	100	
SVAAATLGFAY	1262		11	14	100	
TIMAKNEYF	2590		9	11	79	
TLHGPTPLY	1622		10	11	79	0.0300
TLLFNLLGW	1811		10	12	86	
TTIMAKNEYF	2509		10	11	79	
TIMRSVPF	1208		8	12	86	
TVDFSLDPF	1466		10	12	86	
VIDTLCGF	122		9	12	86	
VLAALAAV	1871		8	12	86	
VLEDGNY	167		8	12	86	
VLVDLQAY	1052		9	11	79	
VMGSSYGF	2639		6	11	79	
VMGSSYGFY	2639		10	11	79	
WMNRRAF	1920		6	14	100	
YSPGQFKEF	2648		9	11	79	
YTNVDDQLYGN	1106		11	11	79	
YVGDLCSIF	276		10	12	86	
	79		2			

Table VIII

HCV A02 Super Motif with Blinding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
93	13	1904	AAILRHIV					
86	12	1673	AAIAYAYCL					
79	11	1250	AAQGIVKL					
79	11	1250	AAQGYKVL					
79	11	1250	AAQGYKVL					
79	11	147	AARALAHGV					
79	11	147	AARALAHGVY					
100	14	1264	AATLFGFGA					
93	13	1264	AATLFGGAYM					
86	12	1187	AAVCTRGV					
79	11	1187	AAVCTRGVA					
79	11	1187	AAVCTRGVAKA					
93	13	1690	AISPGAL					
86	12	1890	AISPGALV					
86	12	1890	AISPGALVV					
100	14	150	ALAHGIVHV					
100	14	150	ALAHGIVRVL					
86	12	1717	ALGLDIA					
86	12	609	ALSTGILH					
79	11	1696	ALVYGVVCA					
79	11	1696	ALVYGVVCAA					
79	11	1696	ALVYGVVCAA					
86	12	1602	AQAPPFSWDM					
79	11	1251	AOQYKVL					
79	11	1251	AOQYKVLV					
86	12	77	AQPGPWPPL					
93	13	1265	ATLFGGAYM					
79	11	1354	ATPPGSVT					
79	11	1598	ATVCARAQAA					
100	14	1419	AVAYYRGL					
100	14	1419	AVAYYRGLDV					
79	11	1168	AVCTRGVA					
79	11	1168	AVCTRGVAKA					
100	14	1917	AVGMMNRL					
100	14	1917	AVGMMNRLA					
93	13	1903	CAAILRHIV					
79	11	1530	CAYVETPA					
86	12	2941	CLRGKVPPPL					
86	12	739	CLWMMLLI					
	11	1653	CMASDLEV					
	79							

## ICV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
				0.00067				
79	11	1653	CMSADOLEVV					
79	11	1653	CMSADLEWVT					
79	11	1126	CTGGSSDL					
79	11	1126	CTGGSSDLYL					
78	11	1126	CTGGSOLYLV					
79	11	1190	CTRGAVAKA					
79	11	1190	CTRGAVAKAV					
79	11	555	CTWNNSTGFT					
86	12	1462	CYTQVDFSL					
79	11	1527	DAGCAGVEL					
100	14	1574	DAHFLSQT					
86	12	1855	DILAGYGA					
79	11	1855	DILAGYGGAGV					
79	11	1855	DILAGYAGVA					
86	12	279	DLGGSVFL					
79	11	279	DLGGSVFLV					
86	12	1657	DLEVNTST					
86	12	1657	DLEVNTSTWV					
86	12	1657	DLEVNTSTWL					
93	13	2617	-2617- -DGVVNGEKA- -DGVVCEKMA					
79	11	132	DLMGYPL					
79	11	132	DLMGYPLV					
79	11	132	DLMGYPLVG					
79	11	2412	DLSGSWST					
79	11	2412	DLSGSWSTV					
79	11	1683	DLVNLPA					
79	11	1683	DLVNLPAI					
79	11	1683	DLVNLPAIL					
79	11	1683	DLVNLPAILV					
79	11	1683	DLVNLPAILV					
79	11	2772	DLVNCESA					
86	12	1134	DLVLTTHA					
86	12	1134	DLVLTTHADV					
86	12	321	DMMNNWSPT					
86	12	1339	DQAEATAGA					
86	12	1339	DQAEATGAA					
86	12	1339	DQAEATGAAVL					
86	12	994	DTAACGDI					
86	12	994	DTLTCGFA					
86	12	124	DTLTCGFADL					
86	12	124	DTLTCGFADLM					
93	13	2673	DIRCFDST					

## ICCV ADD Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
93	13	2673	DTRCFDSTV					
93	13	2673	DTRCFDSTV					
86	12	21	DYKPGEGGDI					
86	12	21	DWPGGGGV					
79	11	750	EALENLV					
100	14	2794	EAMTRYSV					
86	12	2237	EAELLWQDEM					
93	13	1377	EIPFGVKA					
93	13	1377	EIPFGVKA	0.0001				
100	14	2814	ELTSCSSSNV	0.0002				
79	11	666	ELSPILLST					
79	11	666	ELSPILLSTT					
86	12	2245	EMGGNITHV					
86	12	1731	EOPKOKAL					
86	12	1731	EOPKOKALGL					
86	12	1731	EOPKOKALGLL					
86	12	1342	ETAGARLV					
86	12	1342	ETAGARLVW					
86	12	1342	ETAGARLVL					
86	12	1342	ETAGARLVL					
86	12	1207	ETTMRSVFT					
86	12	1207	ETTMRSVFT					
86	12	1659	EVNTSTWV	0.0001				
86	12	1659	EVNTSTWV	0.0004				
86	12	1659	EVNTSTWV					
93	13	130	FADLWGYI					
79	11	130	FADLWGYIPL					
79	11	130	FADLWGYIPLV					
100	14	1927	FASRGNHV					
86	12	1927	FASRGNHVSPV					
100	14	1773	FISGIOYL					
100	14	1773	FISGIOYLA	0.1000				
100	14	1773	FISGIOYLAGL					
79	11	1304	FLADGCGGGAA	0.0046				
86	12	177	FLALLSCL					
86	12	177	FLALLSCLT					
93	13	728	FLLDARV					
86	12	1228	FQVAVHLHA					
86	12	1228	FQVAVLHAPT					
79	11	2646	FOYSGQRV					
100	14	2782	FTTEAMTRYSV					
		13	FTGATHDA					
		13	FTGATHDA					

## ICCV\_A02\_Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
93	13	512	FTPSPIVY					
93	13	512	FTPSPIVYGT					
93	13	512	FTPSPIVYGT					
79	11	684	FTTUPALST					
79	11	684	FTTUPALSTGL					
79	11	146	GAARAAHGV					
86	12	992	GADTAACGDI					
86	12	992	GADTAACGDI					
86	12	350	GAHNGVLA					
79	11	1895	GALVGVGV					
79	11	1895	GALVGVGVCA					
79	11	1895	GALVGVGVCAA					
86	12	1345	GARLYVLA					
79	11	1345	GARLYVLAT					
79	11	1345	GARLYVLAT					
79	11	1345	GARLYVLATAT					
100	14	1916	GAVQWNNRL					
100	14	1916	GAVQWNNRL					
100	14	1916	GAVQWNNRL					
100	14	1333	GIGRVDQDA					
100	14	1333	GIGRVDQDAET					
100	14	1776	GIOYLAGL					
100	14	1776	GIOYLAGLST					
100	14	1776	GIOYLAGLSTL					
79	11	1425	GLOVSIPT					
93	13	1552	GLPVOODFL					
79	11	968	GLRQLAVA					
79	11	968	GLRQLAVA					
100	14	1782	GLSTLPGNPA					
79	11	1782	GLSTLPGNPA					
93	13	1569	GLTHIDAHFL					
93	13	26	GCYVCCVYL					
93	13	26	GONGGVILL					
79	11	2063	GTFFPINAYT					
79	11	2063	GTFFPINAYTT					
100	14	1335	GTVDQDAET					
100	14	1335	GTVDQDAETA					
86	12	1869	GYAGALVA					
79	11	1081	GYCWTIVYHGA					

## IICV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0204	A'6802
86	12	1870	GVLALAA					
86	12	1670	GVLALAYCL					
79	11	161	GIVYATGVL	0.0001				
86	12	45	GVRAIAKT					
100	14	2619	GIVRCERK					
100	14	2619	GIVRCERMA					
100	14	2619	GIVRCERML	0.0002				
83	13	154	GIVRAEDGV	0.0001				
	79	11	GIVVCAAL					
100	14	1234	IAPTGSGKST					
100	14	1572	HIDAHFLSQT					
86	12	696	HILHQNDV					
79	11	1719	HILYIEQGM					
93	13	1769	HMWNFESGI					
79	11	698	HQNNNDQYL					
79	11	222	HTPGCIPCV					
86	12	2855	HTPVNSWL					
86	12	2855	HTPVNSWLN					
	79	11	IYGERSEGA					
79	11	1910	HYGRGEAV					
86	12	1910	HVSPTHV					
100	14	1925	IAFASIGNHV					
79	11	1856	ILAGYAGAV					
79	11	1856	ILGGWVAA					
86	12	1816	ILGGWVAOL					
86	12	1816	ILGGWVAOLA					
86	12	1331	ILGIGIVL					
86	12	1331	ILGIGTVLDQA					
93	13	1891	ILSPGALVY					
93	13	1891	ILSPGALVV					
93	13	1891	ILSPGAIVGV					
79	11	2591	IMARNEVFCV					
100	14	1777	IOYLAGLST					
100	14	1777	ITVSESENKV					
86	12	2250	ITVSEENRIV					
86	12	2250	ITSCSSSNV					
100	14	2816	ITSCSSNWSV					
100	14	2816	ITSCSSNSVA					
86	12	969	ITWGADTA					
86	12	569	ITWGADTAA					

## ILCV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802.
79	11	1296	ITSTYGGFL					
79	11	1296	ITSTYGGFLA					
79	11	2613	IVFDGLGV					
79	11	2613	IVFDGLGVAV	0.0016				
93	13	30	IVGAVYLL					
86	12	1736	KALGILLOT					
86	12	1736	KALGILLOTAA					
86	12	2625	KMAYDVV					
86	12	1734	KOKALGIL					
86	12	1734	KOKALGILLOT					
86	12	1734	KOKALGILLOTAA					
86	12	121	KVDTLTCGFA					
86	14	1255	KVLVLPNSV	0.0048				
100	14	1255	KVLVLPNSVAA					
100	14	1255	KVPAVAYAA					
79	11	1244	LAALAYCL	0.0011				
86	12	1872	LADGGGGGA					
79	11	1305	LAEOFKOKA					
86	12	1729	LAEOFKOKAL					
86	12	1729	LAGYGAGV					
79	11	1857	LAGYGAGVA					
79	11	1857	LAGYGAGVAGA					
79	11	1857	LAGYGAGVAGA					
100	14	151	LAHGIVAVL					
86	12	179	LALLSCLT					
79	11	972	LAVVIEPV					
100	14	1924	LIAFASRGNHHV	0.0004				
100	14	2615	LITSCSSNVSV					
79	11	2612	LVFPDLGV	0.0002				
79	11	2612	LVFPDLGVAV					
86	12	178	LLALLSCL					
86	12	178	LLALLSCLT					
86	12	178	LLFLILADA					
93	13	726	LLFLILADHIV					
86	12	1812	LFNRGGW					
86	12	1812	LFNRGGWVA					
93	13	729	LLLADARY					
93	13	1887	LLPAAILSPGA	0.0061				
93	13	1887	LLPAAILSPGAL					
83	13	36	LLPARIGSPFL	0.0025				
83	13	36	LLPARIGSPGV					

## HCV\_A02\_Super Motif with Blinding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
86	12	2240	LLWROEMGGNI					
93	13	1629	LLYRH GAV					
79	11	133	LMGMYFLV					
79	11	133	LMGYVPLVGA					
86	12	2761	LOQCTMVLV					
86	12	126	LTCGFADLM					
86	12	126	LTCGFADLM					
100	14	2160	LTDPSHIT					
100	14	2160	LTOPSHITA					
86	12	1052	LIGRDKNOV					
93	13	1570	LTHIDAHFL					
93	13	2176	LTSMLTGTSHI					
79	11	2739	LTTSGCGNT					
79	11	2738	LTTSGCGNTL					
79	11	2738	LTVAYQATV					
86	12	1591	LVAYQATYGA					
86	12	1591	LVAYQATYGA					
79	11	1853	LVDIRAGYGA					
86	12	1867	LVGGVLA					
86	12	1867	LVGGVLA					
86	12	1667	LVGGVLA					
86	12	1667	LVGGVLA					
100	14	1257	LVINPNSVA					
100	14	1257	LVINPNSVAAT					
100	14	1257	LVINPNSVAATL					
79	11	1684	LVNLPAI					
79	11	1884	LYRHADV					
86	12	1137	LYRHADVI					
79	11	1137	LYRHADVIPY					
79	11	1897	LYVGNCVA					
79	11	1897	LVGGVCAAA					
79	11	1897	LVGGVCAAA					
79	11	1637	LVGNCVAAL					
79	11	1137	LVVICESA					
79	11	2773	LVVLATAT					
86	12	1348	MANNVEFCV					
86	12	2592	MLTDPSH					
100	14	2179	MLTDPSHIT					
100	14	2179	MLTDPSHITA					
100	14	2179	MMNNWSPT					
93	13	322						

## HCV A92 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0205	A'6802
93	13	1418	NAVAYFHGL					
93	13	1418	NAVAYFHGLDV					
86	12	2068	NAYTGPCT					
86	12	1815	NILGGWVA					
66	12	1815	NILGGWVAAGL					
66	12	1815	NIRTGIVFT					
93	13	1282	NIRTGIVFTI					
79	11	1282	NIRTGIVFTIT					
79	11	1282	NIRTGIVFTTT					
66	12	2249	NITRVESENKV					
86	12	700	NIVDVOYL					
86	12	116	NLGKVITC					
86	12	116	NLGKVITCL					
86	12	116	NLGKVITLT					
93	13	1886	NLLPAISPGAG					
86	12	2239	NLWRCDEM					
93	13	166	NLPGCSESI					
93	13	166	NLPGCSESFIL					
86	12	1460	NTCVTQTY					
93	13	416	NTNGSMWH					
86	12	14	NTNRPQDVA					
93	13	1889	PALSGVA					
93	13	1889	PALSGVAL					
86	12	1889	PAILSPGALV					
93	13	1889	PAILSPGALW					
86	12	698	PALSTGLI					
86	12	698	PALSTGLHL					
79	11	2609	PARLWFPDL					
79	11	2066	PINAYTTGPCT					
79	11	1295	PITYSTYGFLL					
93	13	2403	PLEGEPPDQL					
79	11	143	PLGGAARRA					
79	11	143	PLGGAARRAL					
79	11	143	PLGGAATALA					
93	13	1628	PLLYRLGAV					
93	13	1628	PLLYRLGAVV					
79	11	2667	PMGFSTYD					
79	11	2807	POPEYDEL					
79	11	2807	POPEYDEU					
79	11	2807	POPEYDLEL					
93	13	7	PORKYRANT					

## HCV-A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A*0201	A*0202	A*0203	A*0206	A*6802
86	12	109	PTDPRARSNL					
79	11	1473	PTFTIETT					
79	11	1473	PTFTIETT					
100	14	1236	PTGS/ST					
93	13	1236	PTGSGKSTKV					
86	12	1936	PTHYV/PESDA					
86	12	1936	PTHYV/PESDA					
79	11	1821	PTLHGQPTPL					
79	11	1621	PTLHGQPTPL					
79	11	2070	PTLWARMIL					
79	11	2870	PTLWARMILM					
79	11	2870	PTLWARMILMT					
79	11	2870	PTPLYLAL					
100	14	1628	PTPLYLALGA					
93	13	1626	PTPLYLALGA					
93	13	1626	PVNSWLGNI	0.0001				
100	14	2657	PVNSWLGNI	0.0001				
100	14	2657	PVNSWLGNI	0.0001				
66	12	2657	PVNSWLGNI	0.0001				
79	11	2318	PVWHGCP					
93	13	508	PVYCFTPSPV					
93	13	508	PVYCFTPSPV					
86	12	1340	DAETAGARBL					
86	12	1340	DAETAGARLV					
86	12	1340	DAETAGARLV					
86	12	1603	QAPPFSWQDM					
93	13	1595	QATVCAAA					
79	11	1595	QATVCAAA					
93	13	29	QIVGGVYL					
93	13	29	QIVGGVYL	0.0015				
86	12	338	OLLAQDA					
86	12	2184	OLPCEPEPDV					
79	11	2210	QLSAPSLSKA					
79	11	2210	QLSAPSLSKA					
86	12	1465	QTVQFSDLT					
86	12	1229	QV AHLHAPT					
86	12	1166	RAAVCTRGVY					
79	11	1166	RAAVCTRGVY					
100	14	1449	RALAHGVAV	0.0001				
100	14	1449	RALAHGVAV	0.0001				
86	12	2733	RASGVLT					
	11	43	RIGVRAATKT					

## UCY A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
78	11	2916	RLHGLSASL	0.0280	0.0056	0.0180	0.0002	0.0032
79	11	2611	RLVERPDL	0.0690	0.0110	1.0000	0.0100	0.0050
79	11	2611	RLVFDQGV					
79	11	1616	RLKPTLHPT					
			RLLAPITA					
86	12	1029	RLVVLATA					
86	12	1347	RLVVLATAT					
86	12	1347	RLWHYPCT					
100	14	619	RMANODMM					
86	12	317	RMYVGGAWEHL					
93	13	635	ROEMGCVN					
86	12	2243	ROEMGGNT					
88	12	2243	ROEAGGNTFV					
86	12	2243	RTGIVATIT					
79	11	1284	RTGIVATIT					
79	11	1284	RVCEKML					
100	14	2621	RVCEKMLHDV					
86	12	2621	RVESENKV					
86	12	2252	RVESENKV					
79	11	2100	RVGDRM					
86	12	1556	RVLEDGVNVA					
86	12	1556	RVLEDGVNVA					
86	12	2833	RVYMLTRDPT					
79	11	1655	SADLEWTT					
79	11	1655	SADLEWTT					
79	11	2212	SAPSLKAT					
79	11	2212	SAPSLKATCT					
93	13	2207	SASQLSASL					
100	14	175	SIFILLALL					
86	12	175	SIFLLALLSCL					
100	14	1470	SLDPTFTI					
86	12	1470	SLDPTFTIET					
79	11	1470	SLHSYSPGEI					
79	11	2926	SLTGRGCGVG					
86	12	1051	SMLTDPSH					
100	14	2178	SMLTDPSHIT					
100	14	2178	SMLTDPSHITA					
86	12	2163	SOLCEPERDV					
93	13	2209	SOLSAPSL					
79	11	2209	SOLSAPSLKA					
79	11	2209	SOLSAPSLKA					

## UCV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0204	A'6802
9.3	13	56	SOPGRGRGPI					
8.6	12	1242	STKVPAAYA					
7.9	11	1242	STKVPAAYAA					
10.0	14	1784	STLPGNPA					
7.9	11	1784	STLPGNPAI	0.0007				
7.9	11	2	STNPKPDKT					
8.6	12	1663	STWVLGGVV					
8.6	12	1663	STWVLGGVVL					
8.6	12	1663	STWVLGGVLA					
8.8	12	1299	STYGKFLA					
10.0	14	1282	SVAAILGFGA					
8.6	12	1455	SVIDCNCTCV					
8.6	12	1455	SVIDCNCTCVT					
8.0	12	995	TAQGDII					
8.6	12	1343	TAGARLVLV					
8.6	12	1343	TAGARLVLV					
8.6	12	1343	TAGARLVLVA					
7.9	11	1343	TAGARLVLAT					
7.9	11	2852	TARHPPVNSVL					
7.9	11	2590	TIMAKNEV					
9.3	13	1266	TLGFEGAYM					
8.6	12	1266	TLGFEGAYMSKA					
7.8	11	622	TLHGPPFL					
7.9	11	1822	TLHGPPFLL					
8.6	12	811	TLPALSTGL					
7.9	11	686	TLPALSTGL	0.0003				
7.9	11	686	TLPALSTGL	0.0004				
7.9	11	1785	TLPGNPAI					
8.6	12	125	TLICGFADL					
8.6	12	125	TLICGFADL	0.0003				
7.9	11	2871	TLWARMILM					
7.9	11	2871	TLWARMILM					
8.6	12	1464	TQVDESDLQPT					
7.9	11	2589	TTMAKNEV					
7.9	11	885	TTLPALSTGL					
7.9	11	685	TTLPALSTGL					
7.9	11	685	TTMRSPVFT					
8.6	12	1208	TTSCGNIL					
7.8	11	2738						

## HCV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
79	11	2739	TTSCGNLT					
79	11	1597	TVCARADA					
86	12	1466	TVDFSLOPT					
86	12	1466	TVDFSLOPT					
100	14	1336	TVLDDAET					
100	14	1336	TVLDQNETAGA					
86	12	1336	TVLDQNETAGA					
100	14	1263	VAATLGFGA					
93	13	1263	VAATLGFGAYM					
88	12	1230	VAHLHAPT					
86	12	1440	VATDNLMT					
86	12	1592	VAYQATVCA					
79	11	1592	VAYQATVCA					
100	14	1420	VAYYRGDGV					
100	14	1420	VAYYRGDGV					
86	12	1456	VIDCNCV					
86	12	1456	VIDGNTCV					
86	12	1456	VIDGNTCV					
86	12	122	VIDLITGFA					
86	12	1671	VLAALAYCL					
93	13	1521	VLCECYDA					
79	11	1521	VLCECYDAAGCA					
100	14	1337	VLDQAEATA					
86	12	1337	VLDQAEATA					
86	12	157	VLEDGVNVA					
86	12	157	VLEDGVNVA					
100	14	1258	VLNPSSAA					
100	14	1258	VLNPSSAA					
79	11	2737	VLTTSGNT					
79	11	2737	VLTTSGNT					
100	14	2737	VLTTSGNTLT					
100	14	1256	VLVDILAGYGA					
79	11	1652	VLVGGVLA					
86	12	1666	VLVGGVLA					
86	12	1666	VLVGGVLA					
86	12	1666	VLVGGVLA					
100	14	1256	VLVLPNSV					
100	14	1256	VLVLPNSV					
100	14	1256	VLVLPNSV					
79	11	2600	VOPEKSGKPA					

## HCV A02 Super Motif with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0208	A'6802
				A'0201	A'0202	A'0203	A'0208	A'6802
100	14	1918	VOWMNLFLV					
100	14	1918	VOWMNLFLV					
100	14	1918	VOWMNLFLFA					
86	12	1463	VTOVDFSL					
79	11	1138	VTRHADVI					
79	11	1138	VTRHADVPV					
86	12	1661	VTSTMVLV					
86	12	1661	VTSTMVLVGGV					
79	11	1439	WVATDLM					
79	11	1439	WVATDLMT					
79	11	1901	WVCAAILTRHV					
79	11	1998	WVGVCVAAI					
79	11	1998	WVGVCVAAI					
79	11	1898	WVGVCVAAIL					
86	12	1660	WVTSTWVL					
86	12	1660	WVTSTWVLV					
86	12	1766	WAKHAWNFA					
86	12	76	WQOPGYRPL					
86	12	2873	WARMILMT					
79	11	2297	WARPDPNPL					
100	14	1920	WMNRLJFA					
79	11	557	WMNSTGFT					
86	12	1665	WVLGGGL					
86	12	1665	WVLGGVLA					
86	12	1665	WVLGGVLAAL					
79	11	1249	YAAGQGYKV					
79	11	1249	YAAGQGYKVL					
79	11	1249	YAAGQGYKVL					
79	11	1249	YIPLVGAPI					
79	11	136	YLAGLSTL					
100	14	1779	YLGSSGGPL					
86	12	1165	YLGSSGGPL					
86	12	1165	YLTFRHADVI					
93	13	35	YLTFRHADVI					
79	11	2836	YLTFRHADVI					
86	12	1590	YLVAYQAT					
86	12	1590	YLVAYQATV					
86	12	1590	YLVAYQATVCA					
86	12	1138	YLVTRHADV					
79	11	1136	YLVTRHADV					
79	13	1594	YQATVCAH					
93								

## MCV\_A02\_Super Moth with Binding Information

Conservancy	Freq.	Position	Sequence	A'0201	A'0202	A'0203	A'0206	A'6802
79	11	1594	YQATVCARACIA					
79	11	1106	YTNQQL					
79	11	1106	YTNQQLV					
86	12	276	YVGDLCSV	0.0018				
86	12	276	YVGDLGSVFL					
93	13	637	YVGAEPL	0.0008				
86	12	1939	YFESDAA					
86	12	1939	YFESDAAA					
86	12	1939	YFESDAAKV					
			555					

Table IX  
ICY A03 Super Motif (With Binding Information)

Conservancy	Freq.	Position	Sequence	A'0301	A'1101	A'3101	A'3301	A'8801
86	12	647	AACWWTGER	0.0003	0.0140	0.0450	0.0055	0.0018
79	11	147	AARALAHGVRA					
79	11	187	AAVCTRGVAK					
78	11	2206	ASCOLSAPSJK					
85	12	1265	ATLGFGAYMSK					
78	11	49	ATRKTSER					
79	11	188	AVCTRGVAK					
86	12	2941	CLRLGIPPLR					
79	11	555	CTWMNSTGFTK					
79	11	2599	COPENGGR					
79	11	2899	COPERKGSK					
100	14	1574	DANHLSQTK					
93	13	2617	DGVYRCEK					
79	11	1143	DIVPVRHR					
86	12	2245	EMGGNTT					
86	12	2946	EVFOVOPK					
100	14	728	FLLADAT					
79	11	146	GAAHALAHGVRA					
100	14	1916	GAVOMMNR					
79	11	3037	GIVLPLNR					
79	11	1004	GLPVSAIR					
86	12	1131	GSSDLILVTR					
86	12	1683	QVAGALVAK					
79	11	3035	GVATLPLNR					
79	11	45	GVTRATKTSER					
79	11	1900	GVVCAAILR					
79	11	1900	GVVCAILR					
93	13	33	GVYLPHPR					
93	13	33	GVYLPHRPR					
79	11	1141	HADIVPVH					
79	11	1141	HADIVPVH					
79	11	1141	HADIVPVH					
100	14	1234	HAPTGSKK					
93	13	1234	HAPTGSKK					
100	14	1572	HIAHFLSOTK					
86	12	1232	HJAPHTGSKK					
100	14	1395	HJIFCHSK					
100	14	1395	HJIFCHSKK					
100	14	1395	HJIFCHSKK					
79	11	2120	HYSIPEINR					
79	11	2222	HTPGCVTCVR					
86	12	2250	ITRVESENK					
86	12	1298	ITSTYGR					
79	11	2813	IPFDQVRA					
93	13	30	IVGGYVLLPR					
93	13	30	IVGGYVLLPR					
86	12	2944	KLGVPPLR					
86	12	10	KTKRANTNR					
86	12	10	KTKRANTNR					
93	13	51	KTSERSOPR					
86	12	51	KTSERSOPR					
88	12	1729	LAEQFKK					

## ICV/AB1 Suncer Motif (With Binding Information)

Conservancy	Freq.	Position	Sequence	A'·0301	A'·1101	A'·3101	A'·3301	A'·8801
86	12	2235	LEANLLWR	0.0008	0.0005	0.0016	0.0088	0.0008
100	14	1396	LECHSKK	0.5400	0.1800	0.0071	0.0012	0.0240
100	14	1398	LECHSKKK	0.0003				
79	11	2612	UVFFDOLGVR					
100	14	726	LLFLLLADAR					
93	13	36	LLPRIGPR					
86	12	97	LLSPGCSR					
79	11	1591	LVATQATCVR					
79	11	1	MSTNPKPQR					
79	11	1	MSTNPKPQRK					
86	12	2249	NITVSEENK	0.0010	0.0062	0.0007		
79	11	14	NITRPPDK	0.0010				
79	11	14	PITYSTYK					
79	11	1295	PMGFSYDTH					
79	11	2667	PSPVNGTIDR					
93	13	514	PSWDQMK					
79	11	1607	PTDPRRNSR	0.0008	0.0005	0.0001	0.0008	0.0002
86	12	108	PTGSKSTK	0.0002	0.0002	0.0005		
93	13	1236	PVAVGTTOR	0.0008				
86	12	1340	QAEATAGAH					
93	13	29	QVGGVYLPK					
86	12	289	QLFTESPR	0.7500	0.0330	0.0290	0.0077	3.1000
79	11	289	QLFTESPR					
79	11	2210	QSSAFSLK					
79	11	1106	RAAVCTRGVAK					
100	14	149	FAALAQVR					
79	11	47	RATRKTSE					
79	11	43	RGVYRATR					
79	11	43	RGVYRATR					
100	14	1923	RLIAFASR					
79	11	2611	RLVFFDOLGVR					
100	14	636	TMVNGVNEIR					
93	13	55	TSFPTGPR					
79	11	2207	SASQSLAPSILK					
86	12	1132	SDLYLVR	0.0003	0.0044			
79	11	2	STNPKPQR					
79	11	2	STNPKPQRK					
79	11	2	STNPKPQRK					
86	12	1266	TLFGAYMSK	0.0010	0.0610	0.0005	0.0013	0.0009
79	11	1622	TLIGFTPLYR					
93	13	62	TSERSOPH					
86	12	52	TSERSOPHGR	0.0003	0.0001			
86	12	52	TSERSOPHGR					
79	11	1050	TSLTGRDK					
86	12	1684	VAGALVAFK	0.2400	0.8900	0.0048	0.0025	0.0310
79	11	1592	VAYGATVCAAR	0.0005	0.0038	0.0680	0.0720	0.0280
86	12	1337	VLDQAEFTAGAR					
79	11	1338	VTHHADMPVVR					
79	11	1901	WYCAAILR					
79	11	1888	WGVVYCAAILR					
93	13	517	WVGVTGDR					

## HCY A01 Super Motif (With Blending Information)

Conservancy	Freq.	Position	Sequence	A'0301	A'1101	A'3101	A'3301	A'6801
86	12	93	WAGWLSPR	0.0008	0.0005			
86	12	86	WLLSPRGSR					
100	14	1920	WMNRLAFAASR	0.0530	0.0610	0.0014	0.0420	0.0056
79	11	557	WMNSTGFTK	0.0054	0.0005			
93	13	35	YLPRRGPR					
79	11	2930	YSPGENRA					
100	14	637	YGGVDRR					
86	12	1939	YVPESDAANF	0.0003	0.0001			
		112						

Table X

## HCV\_A24\_Super\_Motif\_With\_Binding\_Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*2401
AIISGAGL	1890	8	1.3	93	
ALAHGIVRL	150	9	1.4	100	
ALSTGIVHL	689	9	1.2	86	
ALVVGIVCAA	1895	11	1.1	79	
ATGNLPGCSF	165	10	1.3	93	
ATLGFGAY	1265	6	1.2	100	
ATLGFGAYM	1265	9	1.3	93	
AVAYYRGL	1419	8	1.4	100	
AVDMNIRL	1917	8	1.4	100	
AVQWMNRL	1917	9	1.4	100	
AVQWMNRLAF	1917	11	1.4	100	
AWDMMMMW	319	8	1.2	86	
AYAAQGYKVL	1248	10	1.1	79	0.0009
AYYRGDLSVI	1421	11	1.4	100	
CLRKGVPP	2941	10	1.2	86	
CLWMMILLI	739	6	1.2	86	
CTCGSSPL	1128	8	1.1	79	
CTCGSSDLV	1128	9	1.1	79	
CTCGSSDLYL	1128	10	1.1	79	
CTRGVAKAVDF	1190	11	1.1	79	
CTNMNSTGF	555	9	1.1	79	
CVOTVDF	1462	8	1.2	86	
CVOTVDFSL	1462	10	1.2	86	
CYDAGCAW	1525	8	1.1	79	
CYDAGCAWY	1525	9	1.1	79	
CYDAGCAYWYEL	1525	11	1.1	79	
DFSLDPITF	1468	8	1.4	100	
DFSLDPITI	1468	10	1.4	100	
DLGGSYFL	279	8	1.2	86	
DLEVNTSTWV	1657	9	1.2	86	
DLEVNTSTWVL	1657	11	1.2	86	
DLGVACGEAM	2617	10	1.3	93	
DLIMGYIPL	132	8	1.1	79	
DLVNLIPAI	1863	9	1.1	79	
DLVNLIPAI	1883	10	1.1	79	
DTAACGDI	994	8	1.2	86	
DTAACGDI	994	9	1.2	86	
DTLTGCFADL	124	10	1.2	86	
DTLTGCFADLM	124	11	1.2	86	
DYKPGSSQI	21	10	1.2	86	
DYPYALMHY	615	9	1.4	100	
EIPFGKAI	1377	9	1.3	93	
ETAGARLVL	1342	10	1.2	86	
ETTMSPVF	1207	9	1.2	86	
EVNTSTWV	1659	9	1.2	86	

## HCV A24 Super Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'2401
FISGQML	1773	8	1.4	100	
FISGQVLAGL	1773	11	1.4	14	
FLLAISCL	177	9	1.2	86	
FTEAMFRY	2792	8	1.4	100	
FIGLTHIDAHF	1567	11	1.3	93	
FTFLPALSTGL	684	11	1.1	79	
FWAKHMMNF	1765	9	1.2	86	
FWAAGHMMNF	1765	10	1.2	86	
GFADLNGY	129	8	1.3	93	
GFADLNGYI	129	9	1.3	93	
GFADLNGYPL	129	11	1.1	79	
GESYDTDF	2689	9	1.1	79	
GIONLAGL	1776	8	1.4	100	
GIONLAGLSTL	1776	11	1.4	100	
GUPVODHIL	1652	9	1.3	93	
GUPVODHLEF	1552	11	1.2	86	
GSAFSLHSY	2921	10	1.1	79	
GLSTLPGNPA	1782	11	1.1	79	
GLTHIDAHF	1569	9	1.3	93	
GLTHIDAHFL	1569	10	1.0	13	
GTFPINAY	2063	8	1.1	79	
GVAGALVAF	1863	9	1.2	86	
GVAKAVDF	1193	8	1.1	79	
GVALARAAV	1670	9	1.2	86	
GVLAALAAVCL	1670	11	1.2	86	
GVNAYAIGLN	161	9	1.1	79	
GVRCVCEOM	2619	8	1.4	100	
GVRCVCEML	2619	10	1.4	100	
GVRCVCEMLY	2619	11	1.4	100	
GVRLMEDGNY	154	11	1.2	86	
GWGRAIL	1800	8	1.1	79	
GWRLLAPI	1027	8	1.1	79	
GWRLLAPITAY	1027	11	1.1	79	
GYGRAGAVAGL	1859	10	1.2	86	
GYPLVQAPL	135	10	1.1	79	
GYFRCRASGL	2728	11	1.2	86	
HJ-HONINDOY	696	11	1.1	79	
HLPYIEQGM	1719	9	1.1	79	
HMWNFESGI	1769	9	1.3	93	
HMWNFESDQY	1789	11	1.3	93	
HTPVNSMVL	2855	8	1.2	86	
HTPVNSMGLN	2855	11	1.2	86	
HNGPGECAQW	1910	11	1.1	79	
IFILLALSLC	176	10	1.2	86	
ILGEMVVAQI	1816	12	1.2	86	

## HCV\_A24 Super Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A 2401
ILGIGIVL	1331	8	12	86	
IMAKNEEV	2591	8	12	88	
ITYSTYKGF	1296	9	12	86	
ITYSTYKGF	1296	10	11	79	
IVDVCVLY	701	8	12	86	
IVGAVVLL	36	8	13	93	
KFPGGGQI	23	8	13	93	
KVIDILTCDF	121	10	12	88	
LFNLGGW	1B13	8	12	86	
LIEANLLW	2235	8	12	88	
LINTNGSW	414	8	11	79	
LLAISLSCL	170	9	12	86	
LLAPITAY	1030	8	14	00	
LFNLGGW	1812	9	12	88	
LPAPALSPGAL	1987	11	13	93	
LPAPALSPGAL	36	9	13	93	
LLSPRSRSPSW	97	11	11	79	
LLWRCOEGAN	2240	11	12	06	
LTCGFADL	126	8	12	86	
LTCGFADNM	126	9	12	86	
LTCGFADLM	126	11	12	86	
LTHIDAHF	1670	8	13	93	
LTHIDAHFL	1570	9	13	93	
LTSMLTOPSH	2176	11	13	93	
LTSQSGANTL	2738	9	11	79	
LVDNLAGY	1853	8	11	79	
LVGGVLAAL	1687	9	12	86	
LVNIPSVATL	1257	11	14	100	
LVNLPAI	1804	8	11	79	
LVNLPAIL	1884	9	11	79	
LVTRHADVI	1137	9	11	79	
LVVGWCAAI	1897	10	11	79	
LVVGWCAAL	1897	11	11	79	
LWARMILM	2872	8	12	86	
LWARMILMIF	2872	11	12	86	
LWRCGANGA	2241	10	12	86	
LYLYTRHADVI	1135	11	11	79	
MILMTHFF	2876	8	12	86	
MLTOPSHI	2179	8	14	100	
MVNFSFGI	1770	8	14	100	
MVNFSFGY	1770	10	14	100	
MVNFSFGYI	1770	11	14	100	
MVNFSFGYL	838	10	13	93	0.0270
NFISGQY	1772	8	14	100	
NFISGQYL	1772	9	14	100	0.0170

## HCV\_A24\_Super\_Motif\_With\_Binding\_Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A <sup>2401</sup>
NIIGGWAACQ	1815	11	12	86	
NIRIGVRTI	1282	9	11	79	
NIVDVCYL	700	8	12	86	
NIVDVQYLY	700	9	12	86	
NIGRIVDTL	118	9	12	86	0.0001
NIIVDRCM	2253	8	12	86	
NIPGCSFSI	168	9	13	93	
NIPGCSFSIF	188	10	13	93	
NIPGCSFSIFL	168	11	13	93	
NTCVTGIVDF	1460	10	12	86	
NTNGSMHI	416	8	13	93	
NTNRFPODFK	14	11	11	79	
NWODGLNW	1108	9	11	79	
NWFGCTTAA	551	8	12	86	
PITYSTYGF	1295	10	11	79	
PITYSTYKFL	1295	11	11	79	
PLEGERQDPL	2403	11	13	93	
PLGAAARAL	143	9	11	79	
PMGFSYDTRCF	2667	11	11	79	
PTDPRRSANL	109	11	12	86	
PTLHGPTIL	1621	9	11	79	
PTLHGPTPLL	1621	10	11	79	
PTLHGPTPLY	1621	11	11	79	
PTLWARMIL	2870	8	11	79	
PTLWARMIL	2870	9	11	79	
PTPLYRL	2870	10	11	79	
PVCOCHLEF	1626	8	14	100	
PVCOCHLEF	1554	9	12	86	
PVCOCHLEF	1554	10	12	86	
PVNSALGN	2867	9	14	100	
PVNSALGN	2857	10	14	100	
PVNSALGNII	2857	11	12	86	
PVNSALGNII	2318	9	11	79	
QFQKQAGL	1732	9	12	86	
QFQKQAGL	1732	10	12	86	
QIGGGVYL	29	8	13	93	
QIVDFSLDPTF	1465	9	13	93	
QIVDFSLDPTF	1919	9	14	100	
QIVDFSLDPTF	1778	9	14	100	
QIVDFSLDPTF	2847	10	11	79	0.0180
QIVDFSLDPTF	2647	11	11	79	0.0180
QIVDFSLDPTF	2918	8	12	86	
RLHGLSAFSL	2918	10	11	79	0.0001
RLHGLSAFSL	2611	8	11	79	

## HCV A24 Super Motif With Binding Information

Sequence	Position	Peptide No.	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*2401
RLLAPITAY	1029		9	12	66	
RMAYNDINMM	317		8	12	66	
RMANDDMMNNW	317		10	12	66	
RMLMLTHF	2875		8	12	66	
RMLMLTHFF	2875		9	12	66	
RAYVGSVVEHFL	635		11	13	93	
RCYCEKML	2821		8	14	100	
RCYCEKMLW	2821		9	14	100	
RLEDGVNY	156		9	12	86	
SFSFELLAL	173		9	12	86	
SFSFELLAL	173		10	14	100	0.0041
SIFLALL	175		8	14	100	
SIFLALLSCL	175		11	12	86	
SLDPFIFTI	1470		8	14	100	
SJFSYSVSPGEI	2928		10	14	100	
SMALDPSH	2178		9	11	79	
STIKPAAV	1242		8	14	100	
STLPGNIPAN	1784		9	12	86	
STWVLVGGVL	1863		10	12	79	
SVAAATLG	1262		8	14	100	
SVAAATLGAY	1262		11	14	100	
SWDQMMKQL	1608		9	11	79	
SWLGNIM	2860		8	12	86	
SYLKGSQQGPL	1164		11	12	86	
TIMAKNEVF	2590		9	11	79	
TLGGAYM	1266		8	13	93	
THQPTPL	1622		8	11	79	
THQPTPL	1622		9	11	79	
TIGPTPL	1622		10	11	79	
TLFNIGGN	1811		10	12	60	
TLPALSTGL	686		9	11	79	
TIPALSTGL	686		10	11	79	
TDGPNPAI	1785		8	11	79	
TLTCGFADL	125		9	12	86	
TLTCGFADLM	125		10	12	86	
TLWARMIL	2871		8	11	79	
TLWARMILM	2871		9	11	79	
TTIMAKNEVF	2589		10	11	79	
TTLPALSTGL	685		10	11	79	
TTLPALSTGLI	685		11	11	79	
TTMRSPVF	1208		8	12	86	
TTSGCNTL	2739		6	11	79	
TVDSLDPTF	1486		10	12	86	
TVMNISTGF	556		8	11	79	
TVWVNGGM	1864		9	12	86	

## HCV\_A24\_Super\_Motif\_With\_Binding\_Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*2401
TYSTYGF	1297	8	1.3	93	
TYSTYGRK	1297	9	1.2	86	0.0230
VFTGLTH	1566	8	1.3	93	
VIDLTCGF	122	9	1.2	86	
VLAALAAAY	1671	8	1.2	86	
VLAALAYCL	1671	10	1.2	86	0.0070
VLEDGAVV	157	8	1.2	86	
VLNPSVAVTL	1258	10	1.4	100	
VLTTSGCITL	2737	10	1.1	79	
VLDILAGY	1852	9	1.1	79	
VLGGGVLAAL	1868	10	1.2	86	
WGSSYGF	2639	8	1.1	79	
WGSSYGRF	2639	10	1.1	79	
VTQTVDFSL	1463	9	1.2	86	
VTRHADVI	1138	8	1.1	79	
WATDALM	1439	8	1.1	79	
WGIVVCAAJ	1898	9	1.1	79	
WGIVVCAAL	1899	10	1.1	79	
WTSTIWL	1860	8	1.2	86	
WLLP-FRGPRL	34	11	1.3	93	0.0016
WINWRLAF	1920	8	1.4	100	
WMLVGGW	1665	8	1.2	86	
WVLVGGVAAAL	1665	11	1.2	86	
YIPLYGAPL	136	9	1.1	79	
YLAGLSTL	1779	8	1.4	100	
YLGSSGGPQ	1165	10	1.2	86	
YLGSSGGPQL	1165	11	1.2	86	
YLPLTRGPRL	35	10	1.3	93	0.0001
YLVTRHADVI	1136	10	1.1	79	
YTMDDC	1106	6	0	11	
YTNDDQDGVW	1106	10	1.1	79	
YVGLOGSIF	276	10	1.2	86	
YVGLOGSFL	276	11	1.2	86	
YVGVEFRL	637	9	1.3	93	
YRGGLDVSIV	1422	10	1.4	100	
	260	3			

Table XI  
ICV B07 Super Motif (with Binding Information)

Conservancy	Pos.	Sequence	B'0702	B'3501	B'5101	B'5301	B'5401
86	12	1804	APPSSWQMW	0.0028	0.0002	0.0001	0.0002
79	11	1804	APPSSWQMW	0.0001	0.0001	0.0006	0.0003
93	13	1235	APTSGSRKTV	0.0001	0.0001	0.0012	0.0002
79	11	2869	APTLWARM	0.4300	0.0001	0.0012	0.0023
79	11	2869	APTLWARMIL	0.0160	0.0002	0.0012	0.0001
79	11	2869	APTLWARMILM	0.0009	0.0001	0.0010	0.0003
79	11	2869	OPOLSDGSW	0.0130	0.0001	0.0003	0.0002
79	11	2410	DFARASRN	0.0001	0.0002	0.0002	0.0005
86	12	111	FPOLGIVR	0.0170	0.0002	0.0001	0.0002
79	11	2815	FRGGQAV	0.0001	0.0001	0.0001	0.0002
100	14	24	FRGGQAVGV	0.0001	0.0001	0.0002	0.0002
86	86	12	1912	GPGECAVQW	0.0001	0.0002	0.0001
86	86	12	1912	GPGECAVQW	0.0001	0.0002	0.0001
93	93	41	GPRLGIVR	0.0001	0.0001	0.0001	0.0002
100	14	1025	GPITPLYL	0.0024	0.0002	0.0002	0.0001
93	93	13	1625	GPITPLYLGA	0.0005	0.0005	0.0005
93	93	13	507	GPYNCFTPSV	0.0001	0.0001	0.0002
93	93	13	1378	IPFYGKAI	0.0120	0.0001	0.1200
79	79	11	137	IPLVGAPL	0.4400	0.0032	0.0700
86	86	12	2808	KPARLIVF	0.0150	0.0002	0.0017
79	79	11	2808	KPARLIVFFDL	0.0003	0.0003	0.0002
79	79	11	1820	KPTLHGPTPL	1.4150	0.0001	0.0002
79	79	11	1820	KPTLHGPTPL	0.0021	0.0001	0.0001
93	93	13	1089	LPAILSPGAL	0.0001	0.0001	0.0002
93	93	13	1888	LPAILSPGAL	0.0053	0.0003	0.0036
86	86	12	1888	LPAILSPGAL	0.0003	0.0003	0.0001
100	14	807	LPALSTGCL	0.0020	0.0001	0.0002	0.0005
86	86	12	687	LPALSTGHL	0.0350	0.0002	0.0002
86	86	12	687	LPCEPEPDV	0.0011	0.0001	0.0001
93	93	13	2665	LPGCSFSI	0.0001	0.0002	0.0002
93	93	13	199	LPGCSFSIF	0.0110	0.0360	0.0059
93	93	13	169	LPGCSFSIIL	0.1950	0.0796	0.0550
93	93	13	169	LPGCSFSIIL	0.0022	0.0009	0.0100
93	93	13	37	LPGRGSPRL	0.0007	0.0001	0.0140
93	93	13	37	LPHKGPFLGV	6.5000	0.0001	0.0180
93	93	13	1553	LPVCCDHL	0.1900	0.0001	0.0009
86	86	12	1553	LPVCCDHLF	0.0005	0.0005	0.0002
86	86	12	1553	LPVCCDHLF	0.0001	0.0048	0.0110
86	86	12	1720	LPYIEGSM	0.0001	0.0001	0.0040
100	100	14	1260	NPSVAAATL	0.0111	0.0001	-0.0002
100	100	14	1260	NPSVAAATLGF	0.0001	0.0002	0.0001
86	86	12	1605	PPPSWQMW	0.0003	0.0001	0.0001
79	79	11	1605	PPPSWQMW	0.0001	0.0002	0.0002
79	79	11	1606	PPPSWQMWNN	0.0002	0.0001	0.0002
79	79	11	1606	PPPSWQMWNN	0.0001	0.0001	0.0013
79	79	11	2317	PPVWRHCP	0.0140	0.0001	-0.0002
79	79	11	2601	OPRKCGRPA	0.0011	0.0001	0.0002
79	79	11	2808	OPFQAE	0.0002	0.0001	0.0180
79	79	11	2808	OPYQOLEU	0.0001	0.0002	0.0002
86	86	12	78	QPGRPMFL	0.0006	0.0001	0.0002

## ICV D07 Super Multi (with Binding Information)

Conservancy	Freq.	Position	Sequence	B·0702	B·3501	B·5101	B·5301	B·5401
B6	12	78	OPGYPMPLY	0.0001	0.0011	0.0002	0.0001	0.0002
93	13	57	CPGEGRGPPI	0.2300	0.0002	0.0001	0.0001	0.0002
79	11	2298	HPDTNIPPL	0.0050				
93	93	13	SPGALVVGV	0.0001	0.0002	0.0002	0.1200	0.0002
79	79	11	SPGALVGVW	0.0001	0.0001	0.0016	0.0001	0.0003
79	11	2931	SPGEINTRW	0.0007				
79	11	2931	SPGEINRVA	0.0003	0.0001	0.0001	0.0002	0.0037
79	11	2649	SPGCRVEF	0.0027				
79	11	2649	SPGCRVERL	0.1200	0.0002	0.0002	0.0001	0.0002
79	11	99	SPRGSRPSW	0.3800	0.0002	0.0005	0.0001	0.0002
86	12	1935	SPTHYVPESDA	0.0001				
86	12	1975	TPCGSGWL	0.0028				
79	11	1126	TPCTCGSSDL	0.0005	0.0001	0.0002	0.0001	0.0003
79	11	1126	TPCTCGSSDLY	0.0001				
86	12	2223	TPGCVPCV	0.0001				
93	13	1550	TPGLIPVODDHL	0.0001				
93	93	13	TPLYRLGA	0.0083	0.0001	0.0002	0.2300	0.0110
93	93	13	TPLYRLGAV	0.0120	0.0001	0.0008	0.0001	0.0003
93	13	1627	TPVNSWLGNI	0.0001	0.0001	0.0053		
86	12	2856	TPVNSWLGNI	0.0001	0.0001	0.0006		
86	12	2856	VPESDAAA	0.0022				
86	12	1940	VPESDAAA	0.0001	0.0001	0.0010	0.0001	0.0003
86	12	788	WPPLLLL	0.0021				
616	14	616	YPVRLWHY	0.0001				
100				76				

Table XII HCY B27 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
AKHMMWIFI	1767	8	1.2	86
AKNEVFCV	2593	8	1.2	66
APALAHGV	148	8	1.4	100
DRSELSPL	663	8	1.1	79
EKGGGRPA	2603	8	1.1	79
ERMLALYDV	2624	8	1.2	86
FROKALGL	1733	8	1.2	86
GHRMADM	315	8	1.3	93
GKSTKVP	1240	8	1.2	86
GRIPARLU	2606	8	1.1	79
HRMADM	316	8	1.3	93
KGGRPHL	1390	8	1.1	79
IRIGVRTI	1283	8	1.1	79
KKCDELAA	1403	8	1.4	100
KKKCDDELA	1402	8	1.4	100
LHGPPTPL	1623	8	1.1	79
LHQNNDV	697	8	1.2	86
LRLAVAV	969	8	1.1	79
NHNSPPHY	1932	8	1.2	86
PGRGRPPI	58	8	1.3	93
PGRGRPSW	100	8	1.1	79
PRRSRNL	112	8	1.2	86
RHAQIVP	1140	8	1.1	79
RHTPYNFW	2854	8	1.2	86
RKLGVPP	2943	8	1.2	86
RKPARLIV	2607	8	1.1	79
RKCRASGV	2730	8	1.3	93
RQGFPLGV	39	8	1.3	93
RPPQDKKF	17	8	1.2	86
SKKCDDEL	1401	8	1.4	100
SPM4GIV	116	8	1.2	86
THIDAHFL	1571	8	1.3	93
TKLKLTP	2985	8	1.2	86
TKVPAAYA	1243	8	1.2	86
TRCFDSTV	2674	8	1.4	100
TRGVAKAV	1191	8	1.1	79
VRVCEKMA	2620	8	1.4	100
VRMLEDGV	155	8	1.3	93
YRGDVS	1423	8	1.4	100
YRHTPYNFW	2853	9	1.1	79
ARLWVPPOL	2610	9	1.1	79
ARLWVLTATA	1348	9	1.1	79
ARMLMLTHF	2874	9	1.2	86
ARPDYNPPL	2298	9	1.1	79
DRSELSPL	663	9	1.1	79

## HCV B27 Super Motif

Sequence	Position	Peptide No.	No of Amino Acids	Sequence Frequency	Conservancy (%)
EQKALYDVV	2624		9	12	86
FKCKALGLL	1733		9	12	86
GHRMADWDM	315		9	13	93
GKSTKVPAA	1240		9	12	86
GRKPARLV	2608		9	11	79
HRMADWDM			9	12	86
IKGGERHUF	1390		9	11	86
KKKCDELAA	1402		9	14	100
LHGLSAFSL	2919		9	11	79
LHGFTPPLY	1623		9	11	79
LHSYSPGEI	2927		9	11	79
LKGSSGGPL	1166		9	12	86
LRKLGVPPL	2942		9	12	86
NHYSPTHVV	1932		9	12	86
NHRPODKF	16		9	11	79
PRRGPLGV	38		9	13	93
RHTIPVNSML	2854		9	12	86
RHNGPGEAA	1909		9	11	79
RKPARLIVF	2607		9	11	79
RKRCRASGVL	2730		9	12	86
RSSRNLRV	114		9	12	86
SKKKCDELA	1401		9	14	100
THYVPESDA	1937		9	12	86
TKVPAAYAA	1243		9	11	79
TRHADVIPV	1139		9	11	79
TRVESENKV	2251		9	12	86
WFPGGGQI	22		9	13	93
YRVCERKML	2620		9	14	100
WLLAPITA	1028		9	11	79
WREBGGN	2242		9	12	86
YRGLDVSF	1423		8	14	100
YRRCRASGV	2729		9	13	93
ARALAHGVRY	148		10	14	100
ARAGQPPPSW	1660		10	11	79
ARHTPVNSWL	2853		10	11	79
ARMLMLTHFF	2874		10	12	86
CHSKKKCDEL	1359		10	14	100
DDRSSELSPL	661		10	11	79
DRSELSPUJ	663		10	11	79
EKGGRPARL	2603		10	11	79
FRAAVICTRGV	1185		10	12	86
GHRMADWDM	315		10	12	86
GKSTKVPAY	1240		10	12	86
GRKPARLV	2606		10	11	79
KHAWNFSGI	1768		13	13	93

HCY B27 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy [%]
KKQDELAALKL	1403	10	12	86
LNQINIVDQY	697	10	11	79
LGSSSGSRLL	1168	10	12	86
QKALGLLQTA	1735	10	12	86
PRNGPGEESAV	1909	10	11	79
PRASPRLGWHA	39	10	13	93
PRPHNGPGEAGA	1908	10	11	79
PRHSPENALGKV	113	10	12	86
PRASRNLGKV	114	10	12	86
SHFGYGGAKD	2552	10	12	86
SHRKKCDELAA	1404	10	14	100
THYVPESDAA	1937	10	12	86
TRGVAKAVDF	1191	10	11	79
TRVSEENKAV	2251	10	12	86
VKPGGGGSW	22	10	13	93
VIVCERKMLA	2620	10	14	100
VRKLEDGVNY	155	10	12	86
WRLLAPITAY	1028	10	11	79
YKVVLINPSV	1254	10	14	100
YHRCRASGVL	2729	10	12	86
AHGIVRMEDGV	152	11	13	93
AQHWNWNFISGI	1767	11	12	86
ARLAHGVAVL	148	11	14	100
ARUVFPDLGV	2610	11	11	79
CHSKKKCDELA	1399	11	14	100
DRDRSELSPL	661	11	11	79
EKGGRKPAHJ	2603	11	11	79
FRAAVCTRGVA	1165	11	11	79
GKSTKVKPAAYA	1240	11	12	86
GRIDDTLTCGF	120	11	12	86
HAMAWDNMMWW	316	11	12	86
KKQCDELAALKL	1402	11	12	86
KANTRRPPDV	12	11	12	86
LGIGTFLLYFL	1523	11	11	79
LHQNDNDVQNL	697	11	11	79
LKPTLHGSPTEL	1619	11	11	79
LRPHNGPGEAGA	1907	11	11	79
PRGPFLLVRA	38	11	13	93
PRASRNLGKV	112	11	12	86
PRNGPGEAGA	1908	11	11	79
PRASRNLGKV	113	11	12	86
SRGIGAVSPHY	1829	11	12	86
SRNGKVIDTL	116	11	12	86
THYVPESDAA	1937	11	12	86
VRLEDGVNYA	155	11	12	86

IICV R27 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
YKVLVLMPSVA	1254	11	1.4	100

HCV B58 Super Motif

Table XIII

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
AALRRHV	1804	8	13	93
AAIAAYCL	1673	8	12	86
AAGCYKVL	1250	8	11	79
AAITLGFGAV	1264	8	14	100
AAVCTRAGV	1187	8	12	86
ASIMMAFTA	1793	2	11	79
ASSASSQL	2204	8	14	100
ATLGFGAY	1265	8	14	100
CSFSPL	172	8	14	100
CSG6AYDI	1310	8	12	86
CSRNVSVA	2619	8	14	100
CTCGSSDL	1128	8	11	79
CTRGVAKA	1190	8	11	79
DTAACGDI	994	8	12	86
DTIJCIGFA	124	8	12	86
EAILEMLV	750	8	11	79
EANTRYSA	2794	8	14	100
ESDAAAARV	1942	8	12	86
ETAGARLV	1342	8	12	86
ETTMRSPPV	1207	8	12	86
FADLMGVI	136	8	13	93
FASRGNIV	1927	8	14	100
FSFLLL	174	8	14	100
FSYDTRCF	2670	8	11	79
FTEAMTRF	2792	8	14	100
FTFSPIVW	512	8	13	93
GAGVAGAL	1861	8	12	86
GAHMGVLA	350	8	12	86
GALVGVW	1895	8	11	79
GARLVLVA	1345	8	12	86
GSGKESTKV	1238	8	13	93
GSSDLYLV	1131	8	12	86
GSSGGPL	1168	8	12	86
GSSYGFQY	2841	8	11	79
GTFPINAY	2083	8	11	79
HSYSPGEI	2928	8	11	79
HTPWNNSWL	2855	8	12	86
ISGQYLA	1774	8	14	100
ITSCSSNN	2616	8	14	100
ITWGADTA	989	8	12	86
KSTKVPAA	1241	8	12	86
LAGYGAGV	1857	8	11	79
LAHGVRVL	151	8	14	100
LAVAEPV	972	8	11	79
LSAPSLKA	2211	8	11	79

## HCV B58 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
LSPGALV	1892	8	13	93
LSTGLHL	690	8	12	86
LTCGFADL	126	8	12	86
LTHIDAHF	1570	8	13	93
MSADLEEV	1654	8	11	79
NSWLGNI	2869	8	14	100
NTCVTQTV	1460	8	12	86
NTNGSMWH	416	8	13	93
PAILSPGA	1889	8	13	93
PAIESTGL	688	8	12	86
PILWARMI	2870	8	11	79
PTPLVRL	1626	8	14	100
QATVICARA	1595	8	13	93
PRARPRWFM	3019	8	14	100
ASELSPLL	664	8	11	79
FSRNGLKV	115	9	12	86
SASFSLHSY	2923	6	11	79
SEASQSLSA	2206	8	14	100
STKVPAAY	1242	8	12	86
STLPGNPA	1784	8	14	100
STLPGQAM	2653	8	12	86
STYGKFLA	1289	8	12	86
TAACGSDII	995	0	12	86
TAGARILV	1343	8	12	86
TTMRSFVF	1208	8	12	86
TTSCGHTL	2739	8	11	79
VAGALVAF	1664	8	12	86
VRHADVI	1138	8	11	79
VTSTWVLV	1681	8	12	86
WAKHMMWF	1766	0	12	86
WARYLVM	368	8	14	100
WQPGYFW	76	8	12	86
YAGQGYKV	1249	8	11	79
YSEPLDL	2905	8	11	79
YSTYGKPL	1298	8	12	86
YTMDDQL	1106	8	11	79
AAKLODCTM	2758	9	16	114
AAQGKVKLV	1250	9	11	79
AARALAHGV	147	9	11	79
AAVLGFGAV	1264	9	14	100
AAVCTRGVA	1187	9	11	79
ASQSLSAFL	2208	9	13	93
ATLQFGAYM	1265	9	26	186
ATVCAIAOA	1596	9	11	79
CAAIRRHIV	1903	9	13	93

## HCV B58 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
CANVETPA	1630	9	11	79
CSFSFLA	172	9	14	100
CGGGAYDI	1310	9	12	86
CTGCGSSDLY	1128	9	11	79
CTFGVAKAV	1190	9	11	79
CTWMMNSTGF	555	9	11	79
DIACCAWYEL	1527	9	11	79
DIAAAGGDI	994	9	12	86
DIFCFDFSTV	2673	9	13	93
ETAGARLV	1342	9	12	86
ETTMARSPVF	1207	9	12	86
FSIFLLALL	174	9	14	100
FSLDPFTI	1469	9	14	100
FTGLTHIDA	1567	9	13	93
GAGVAGALV	1851	9	12	86
GALVAFKIM	1866	9	12	86
GALVAFKVM	1866	9	14	100
GAVQMMNRL	1916	9	14	100
HSKKKCDEL	1400	9	14	100
HTPGCVPCV	222	9	11	79
ITWGADTA	989	9	12	86
ITSYTGKF	1296	9	12	86
KALGLLOTA	1736	9	12	86
KSKVVPAY	1241	9	12	86
LAALAAAYCL	1672	9	12	86
LAEQFRKIA	1729	9	12	86
LAGLAYYSM	356	9	14	100
LAGYGAGVA	1657	9	11	79
LSAFSLHSY	2922	9	11	79
LSTLPGMPA	1783	9	14	100
LTCGFAIDL	126	9	24	171
LTFPSHTVA	2180	9	14	100
LTFADKNOV	1052	9	12	86
LTHDAHFL	1570	9	13	93
LTTSCGNTL	2738	9	11	79
MARNEVFCV	2592	9	12	86
MAMDMMMMMW	318	9	12	86
NAVAYYRGL	1418	9	13	93
NSLRLH-FRM	2481	9	14	100
NWLGNIM	2858	9	24	171
NTNFRPQDV	14	9	12	86
PAILSPGAL	1869	9	13	93
PSVAAATLGF	1261	9	14	100
PTLH-QPTPL	1621	9	11	79
PTLWARMIL	2870	9	11	79

## HCV B58 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy [%]
QAEETGAAEL	1340	9	12	8.6
RAAVCTGRV	1186	9	12	8.6
RAIAKIGYRV	148	8	14	10.0
RAQAPPSSW	1601	9	11	7.9
RAYAMDREM	811	9	16	11.4
RSELSPLL	664	2	11	7.9
RSANLGVVI	115	9	12	8.6
SSASASLSA	2205	9	14	10.0
STKVRPAAYA	1242	9	12	8.6
STLPGNPAI	1784	9	11	7.9
STWVLVGGV	1663	9	12	8.6
TAGARILVVL	1343	9	12	8.6
TCSGSNVSV	2817	9	14	10.0
TTIMARKNEV	2589	9	11	7.9
VAATLGFFSA	1263	9	14	10.0
VAGGHHYICM	933	9	14	10.0
VAYQATYCA	1592	9	12	8.6
VAYYRGGLDV	1420	9	14	10.0
VSTLPQAVM	2832	9	12	8.6
VTQINDFSL	1463	9	12	8.6
WAHHMWNF	1766	9	12	8.6
YAOQGYVIL	1249	9	11	7.9
YAPTLWARM	2868	9	14	10.0
YSPGEINAV	2930	9	11	7.9
YSPGEINAV	2846	9	11	7.9
YSTYIGKFLA	1298	9	12	8.6
YTYVIVDQOLV	1106	9	11	7.8
AAQGYYKVLV	1250	10	11	7.9
AATLGGAYAM	1264	10	28	18.6
ASLAVFTEAM	2787	10	12	8.6
ASSASOSLSA	2204	10	14	10.0
ATGNLPGCSF	165	10	13	9.3
CSFSIFLLAL	172	10	14	10.0
CTCGSSDLYL	1128	10	11	7.8
DARVACIWM	733	10	18	12.9
DSWIDGNTCV	1454	10	12	8.6
DTLTCGFADL	124	10	12	8.6
EANLWRCM	2237	10	24	17.1
ETAGARILVVL	1342	10	12	8.6
FADQMGYIPL	139	10	11	7.9
FTEAMTRYSA	2792	10	14	10.0
GAARALAHGV	146	10	11	7.9
GADTAACGDI	992	10	12	8.6
GAGVAGALVA	1861	10	12	8.6
GALVGVVCA	1695	10	11	7.9

## HCV R58 Super-Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
GARLWVLATA	1345	10	1.1	79
GAQWMMARLI	1916	10	1.4	100
GSGKSTRVPA	1238	10	1.2	86
GTIVDQAEATA	1335	10	1.4	100
HSKKKQDEDA	1400	10	1.4	100
IAFASRGNHV	1925	10	1.4	100
ISGICYLAGL	1774	10	1.4	100
ITRVESENKV	2250	10	1.2	86
ITSCSSNSVSV	2818	10	1.4	100
ITYSTYTGKFL	1236	10	1.1	79
KSTKVPAYA	1241	10	1.2	86
LAGGCGSAGA	1305	10	1.1	79
LAEOFKOKAL	1723	10	1.2	86
LALPPRAYAM	806	10	1.2	86
LSPGALVGV	1892	10	1.3	93
LSPRGSRPSW	88	10	1.1	79
LSRARRPRWMF	3017	10	1.4	100
LSTLPGNPA	1763	10	1.1	79
LTHPITKYM	1842	10	1.5	114
NTCVTDTVDF	1460	10	1.2	86
PALSPGALV	1889	10	1.2	86
PALSTGJHL	868	10	1.2	86
PALIVFPDL	2609	10	1.1	79
PSWDDMMMKCL	1607	10	1.1	79
PTSGSGKSTRV	1235	10	1.3	93
PTIYVPESDA	1935	10	1.2	86
PTLHGPTPL	1621	10	1.1	79
PTLWARMILM	2870	10	2.2	157
PTPLVYRLGA	1628	10	1.3	93
QAEFTAGABLY	1340	10	1.2	86
QAPPSPWDM	1603	10	2.4	171
QATVCARAOA	1595	10	1.1	79
RAAKLQDCTM	2757	10	1.6	114
RAAVCTRGVA	1198	10	1.1	79
RALAHGVRL	149	10	1.4	100
SASQSLAPSIL	2207	10	1.3	93
STKVPAAAYAA	1242	10	1.1	79
STWIVLGGVL	1663	10	1.2	86
TAGARLVLVA	1343	10	1.2	86
TAKHTPNWS	2852	10	1.1	79
TCSSNSVSV	2817	10	1.4	100
TSMLTOPSHI	2177	10	1.3	93
TSWWLGGVY	1662	10	1.2	86
TTIMAKNEVF	2589	10	1.1	79
TTLPALSTGL	685	10	1.1	79

## HCV B58 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
VAATLGFGAY	1263	10	1.4	100
VTPGERPSGM	1507	10	1.6	114
VTRHADIVPV	1138	10	1.1	79
WACPGYVWPL	76	10	1.2	86
WARMILNTHF	2873	10	1.2	86
WARPDDYNPPL	2297	12	1.1	79
YAQGQYKVLV	1249	10	1.1	79
YSGENRVA	2930	10	1.1	79
YSGCQEVFL	2648	10	1.1	79
AARLAHGRV	147	11	1.1	79
AASLRVFTEA	2768	11	1.2	88
AAVCTRGVAKA	1187	11	1.1	79
ASHLPHIEQGM	1717	11	1.4	100
ASQIAPSLSKA	2208	11	1.1	79
CARDQAPPSSW	1599	12	1.1	79
CSFSIFLLL	1772	12	1.4	100
CTCGSSOLIVV	1128	12	1.1	79
CTRGIVAKAIDF	1190	12	1.1	79
DARVCACLMMRM	733	12	1.6	114
DTLICGFDLM	124	12	2.4	171
EIAGARLWLA	1342	11	1.2	86
FADLMGYVPLV	130	11	1.1	79
FSL-HSYSPGEE	2925	11	1.1	79
FTQLTHIDAHF	1567	11	1.3	93
FTTPALSTGL	884	11	1.1	79
GADTAACSDII	992	11	1.2	86
GAGVAGALVAF	1861	11	1.2	86
QALVIVQVWCA	1895	11	1.1	79
QAVQMMNRLIA	1918	11	1.4	100
GSKCKSTKVPAA	1238	11	1.2	86
HSRKKCDELAA	1400	11	1.4	100
HSTSPGEIMRV	2928	11	1.1	79
HTPVNSWLNQI	2855	11	1.2	86
ITRVESENKRV	2250	11	1.2	86
ITSCSSENNSVA	2816	11	1.4	100
ITYSTYKPLA	1296	11	1.1	79
KSTKVPAYAA	1241	11	1.1	79
LADGGSGGAGV	1305	11	1.1	79
LAGYGGAGVAGA	1857	11	1.1	79
LNSLRLRHNM	2779	11	1.4	100
LSPGALVGV	1892	11	1.1	79
LTCGFDALMGY	126	12	1.2	86
LTSMLTOPSH	2176	13	1.3	93
NAVAYYRGIDV	1418	13	1.3	93
NTRHPQDNKF	14	11	1.1	79

## HCV\_B58 Super Motif:

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
PAILSPGALVV	1889	11	12	86
PSVATLGGFA	1261	11	14	100
PTDFPFSERNL	109	11	12	86
PTHIVPESDAA	1936	11	12	86
PTLHGPTPLLY	1621	11	11	79
PTPLLYRLGAV	1626	11	13	93
QAEFTAGARLVL	1340	11	12	86
QAPPSPNDMW	1603	11	11	79
QIVDFSLDPTF	1465	11	12	86
PSQPGGRQPI	55	11	13	93
SADIEWVTSIW	1655	11	11	79
SSASQSLSAPSL	2206	11	13	93
SSDLYLVTRAHA	1132	12	12	86
STWVWLGCGA	1663	12	12	86
TARHITPVNSWL	2852	11	11	79
TSLTGDRDNQV	1050	12	12	86
TSTWVWLGCGV	1662	12	12	86
TILPALSTGLI	685	11	11	79
VAAATGEGANAA	1283	26	186	100
VAGALVAFKVM	1664	11	14	86
VAVEPVVFSDM	974	11	12	86
VAYDATVVCARA	1592	11	11	79
VAYFNGSLDVS	1420	11	14	100
VTSWVWLGCGV	1661	12	12	86
WAQPGTGPWPLY	76	12	12	86
WARMILMTHFF	2873	11	11	78
YAQGTYKVALV	1249	11	12	86
YATGALPACSF	164	11	11	79
YTNNDQDQVGW	1106			

## HCV B62 Super Motif

Table XIV

Sequences	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
AILSPGAL	1680	8	13	9.3
ALAHGIVRV	150	8	14	100
ALGLLQTA	1737	8	12	88
APTLWARM	2689	8	11	79
AQAPPSSW	1602	8	12	88
ACGYKVLV	1251	4	11	79
AVAYYRL	1419	8	14	100
AVCTRGIV	1188	8	11	79
AVQWMNRL	1917	14	100	
CLWAMILLI	739	8	12	86
CMSADLEV	1653	8	11	79
CDQHLEFW	1556	8	12	86
CVTOYDF	1462	8	12	86
DILACYGA	1855	8	12	86
DLGGSVLF	279	9	12	86
DLMGYIPL	132	9	11	79
DLYNLPA	1883	8	11	79
DQAEATGAA	1339	8	12	86
EIPFGIKA	1377	13	13	93
ECPIKQAL	1731	8	12	86
EVNTSTWV	1659	8	12	88
FISGIVYL	1773	8	14	100
FPDQGIVV	2615	8	11	79
FPGGGIVV	24	14	100	
FGVAHHLA	1228	8	12	86
GIOYLAGL	1776	8	14	100
GLRDLAVV	968	8	11	79
GPITLEVRA	41	8	13	93
EQVGEAVV	28	6	14	100
GVAGALVA	1863	8	12	86
GVAKAVDF	1193	8	11	79
GVLAALAR	1670	8	12	86
GVVICEM	2619	8	14	100
GVVCAAIL	1800	8	11	79
HVFGGEKA	1910	8	11	79
HVSPTHMV	1930	8	12	86
ILGGWVAA	1816	8	12	86
ILGIGTVL	1331	8	12	86
ILSPGALV	1891	8	13	93
IMAKNEVF	258	8	12	85
IPFGKAI	1378	8	13	93
IPLYGAPL	137	8	11	79
IVDONLY	701	8	12	86
IVPPDGLV	2613	8	11	79
IVGGYLL	30	13	93	

## HCV\_R62\_Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
KMAYLDVV	2625	8	12	66
KPARLIVF	2609	8	12	66
KOKAGLL	1734	8	12	86
KVPAAYAA	1244	8	11	79
LIEANLW	2235	8	12	86
LINTRGSSW	414	8	11	79
LLAISCL	178	8	12	86
LLAPITAY	1030	8	14	100
LLADARV	729	8	13	93
LLYRLGAV	1629	8	13	93
LMGYPLY	133	8	11	79
LPALSTGL	687	8	14	100
LPGCSFSI	168	8	13	93
LPRRAPHL	37	6	13	93
LPVCOCHL	1553	8	13	93
LPYNEQGM	1720	8	12	86
LODCTMVL	2761	8	12	86
LVAYQATV	1691	8	12	86
LVDLILGY	1853	8	11	79
LVGGVLA	1667	8	12	86
LVLNPSVA	1257	8	14	100
LVNLLPAI	1884	8	11	78
LVTRHADV	1137	8	12	86
LVVGVVCA	1897	8	11	79
LYVICESA	2773	8	11	79
MILMLTHFF	2876	8	12	86
MLTDPSHI	2179	8	14	100
NILGGAWA	1815	8	12	86
NIVDQVYL	700	8	12	86
NLWRCM	2239	8	12	86
NPSVAAITL	1260	8	14	100
PLGGAAIA	143	8	11	78
PLTHLGLA	1628	8	13	93
PPPSWIDQM	1605	8	12	86
PPSWDDMW	1606	8	11	79
PVNHGCP	2318	8	11	79
QNGGGYML	29	8	13	93
QLLRIFQIA	336	8	12	86
QPEYDLEL	2808	8	11	78
QPGYPNPPL	78	8	12	86
RLHGLSAF	2918	8	12	86
RLVPPDL	2611	8	11	79
RLAPITAA	1029	8	12	88
RLVVLATA	1347	8	12	88
RMANDMM	3117	12	86	

## HCV B62 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
RAMILTHF	2875	8	12	86
RPDYNPL	2299	9	11	79
ROEMGN	2243	8	12	86
RVCEKMAI	2621	8	14	100
RVESENRY	2252	8	12	86
RVGDRHVV	2100	9	11	79
SIFLLALL	175	8	14	100
SLDPTFTI	1470	8	14	100
SPGENRY	2931	8	11	79
SPGCRVRF	2649	8	11	78
SOLSAPSL	2209	8	13	83
SVAATLGF	1262	8	14	100
TIMAKNEY	2590	8	11	79
TLGFGAYM	1266	8	13	93
TLHGPPTL	1622	8	11	79
TLPGNPAI	1785	8	11	79
TLWARMIL	2871	8	11	79
TPCGSGMIL	1875	8	12	86
TPGCVPCV	223	8	12	86
TOTVDFSL	1464	8	12	86
TVCARAQAA	1597	6	11	79
VIDCNTCV	1456	8	12	86
VLAALAY	1671	6	12	86
VLCCEYDA	1521	8	13	93
VLDQAEITA	1337	8	14	100
VLEDGVNN	157	8	12	86
VLNPSVAA	1253	8	14	100
VLYGGMIA	1686	8	12	86
VLVLNPSV	1258	8	14	100
VMGSSYGF	2639	8	11	79
VPESDAA	1940	8	12	86
VQMANRIL	1918	8	14	100
VVATDALM	1439	8	11	79
VGVVCA	1898	8	12	86
VVTSTWVL	1660	8	14	100
WMANFLAF	1920	8	12	86
WPLLLL	799	8	12	86
WVVGAVL	1665	8	12	86
YLAGLSTL	1779	8	14	100
YPRFLWHY	616	8	14	100
YVPESDAA	1939	8	12	86
AILSPGAVLV	1890	9	12	86
ALAHGVVRL	150	8	14	100
ALSTGVLHL	689	9	12	86
ALVVGVCA	1898	9	11	79

## IICY\_B62 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
APPPSWDOM	1604	9	1.2	86
APTLWARMI	2869	9	1.1	79
AGSGYKVL	1251	9	1.1	79
AGPGYVPL	77	9	1.2	86
AVQWMPRL	1917	9	1.4	100
CMSADLEV	1653	5	1.1	79
DLGSVPLV	279	9	1.1	79
DLIEWTSTW	1657	9	1.2	86
DLIMGYPLV	132	9	1.1	79
DVNLLPAI	1883	9	1.1	79
DLVNCESA	2772	9	1.1	78
DLYLYTRHA	1134	9	1.2	86
CPOLSGSW	2410	9	1.1	79
DPRRRSNL	111	9	1.2	86
EPFPGKAI	1377	9	1.3	93
EMSGENTRV	2245	9	1.2	86
EWSTSTWVL	1658	9	1.2	86
FISGIOYLA	1773	8	1.4	100
FILLSSCL	177	8	1.2	86
FILLDARV	728	9	1.3	93
FONPSGRV	2646	9	1.1	79
GIGTVLDDA	1333	9	1.4	100
GLPVQODHL	1552	9	1.3	93
GLRDIAVAV	968	9	1.1	79
GLTHDAHF	1569	9	1.3	93
GPGEGAVGW	1912	9	1.2	86
GPTFLYRL	1625	9	1.4	100
GQGGGYL	28	9	1.3	93
GIVAGALVAF	1863	9	1.2	86
GVLAAKAY	1670	9	1.2	86
GVNTATGQL	161	9	1.1	79
GVRVCEKMA	2619	9	1.4	100
GYRFLLEDGV	154	8	1.3	93
HJ-HQHNDV	696	9	1.2	86
HLPVIEQGM	1719	9	1.1	79
HMWNFISGI	1769	9	1.3	93
HONNOVY	698	9	1.1	79
HVGREGAV	1910	9	1.1	79
ILASYGAGV	1656	9	1.1	79
ILSPFGALVW	1891	9	1.3	93
KVLYLNPVY	1255	9	1.4	100
LITSCSSNW	2816	9	1.4	100
LWPPDGV	2612	9	1.1	79
LLFLLLADA	726	9	1.4	100
LLFNLQGWW	1812	8	1.2	86

## HCV 362 Super Motif (No binding data)

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
UPRARGPPL	36	9	13	93
LPAILSPGA	1888	8	13	93
LPALSTGLI	687	9	12	86
LPCEPEPDV	2165	9	12	86
LPGCSESF	1693	9	13	93
LVGGVLAAL	1667	9	12	86
LVLNPSVAA	1257	9	14	100
LVNLPLAII	1884	9	11	79
LVTRHADVI	1137	9	11	79
LYVAVCAAA	1697	9	11	79
NILGGWVAA	1815	9	12	86
NIRTGAVR	1282	9	11	79
NIVDQVLY	700	9	12	86
NLGKVDTL	1118	9	12	86
NLPGESFSI	1688	9	13	93
NVQDOLGVAV	1108	9	11	79
PLGGAARAL	1433	8	11	79
PILYRLGAV	1628	9	13	93
PPPSWDDKAV	1605	9	11	79
PPWHGCPPL	2317	9	11	79
POPEYDRL	2807	9	11	78
PVODDHLF	1554	9	12	86
PVNSMIGNI	2857	9	14	100
QVGGVML	29	9	13	93
QLSAPSLSA	2210	9	11	79
QPEYDRL	2808	9	11	79
QPGYFWRLY	78	9	12	86
QPGPQRPPI	57	9	13	93
HLAPITAY	1029	9	12	86
HMILMHNFF	2875	9	12	86
RVCEKMAIY	2621	8	14	100
RVESENKWW	2252	9	12	86
RLLEDGNYI	156	9	12	86
SMLTDFSHI	2178	9	14	100
SPGALVIGV	1893	9	13	93
SPGEMNVA	2931	9	11	79
SPQRVERL	2649	9	11	79
SPRSARPSN	99	9	11	79
SVIDCNTCV	1455	9	12	86
TIMAKNEYF	2590	9	11	79
TLHGTPPLL	1622	9	11	79
TLPALSTGL	686	9	11	79
TLTCGFADL	125	9	12	86
TLWARMILM	2871	9	11	79
TPLYFALGA	1627	9	13	93

## HCV 1062 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
TVLDOAETA	1336	9	1.4	100
VIOTLTGCF	122	9	1.2	86
VLEDGVNVA	157	9	1.2	86
VLVDILAGV	1852	9	1.1	79
VLVGGVLA	1666	24.0075	1.2	86
VLYLNPVVA	1256	24.0072	1.4	100
VOMMNRHIA	1918	9	1.4	100
WGYNVCAAI	1698	9	1.1	79
WTSTWWLV	1660	1.0823	1.2	86
WMNRJAAFA	1920	24.0073	1.4	100
WLYVGGVLA	1665	40.0075	1.2	86
YPLVGVPL	136	1.0817	1.1	79
YLVAYQATV	1590	1.0127	1.2	86
YLVTRHADV	1136	1.0118	1.2	86
YQATVCAARA	1594	9	1.3	93
YVGDLGGSV	276	9	1.2	86
YWGVERFL	637	1.0107	1.3	93
YVPESDAAA	1939	9	1.2	86
AILSPGALVV	1890	24.0101	1.0	86
ALVNGVYCAA	1896	15.0233	1.0	79
APPPSWDQMW	1604	15.0247	1.0	79
APTLWARMIL	2869	1.0107	1.0	79
ACFGYPNPFPLY	77	1.0486	1.0	100
AVAVYRGGLDV	1419	1.0486	1.0	14
AVCTRGVAKA	1188	1.0486	1.0	11
AVQWNNHRLA	1817	1.0510	1.0	14
CLRKLGVPPL	2941	1.0487	1.0	12
CYQTQIVDFSL	1462	1.0487	1.0	12
DILAGYGAGV	1855	1.0485	1.0	11
DLEWTSTWV	1657	1.0490	1.0	12
DLGVRVCEQW	2617	1.0499	1.0	13
OLSDGSMWSTV	2412	1.0499	1.0	11
DLYNLLPAIL	1893	1.0891	1.0	11
DQAEATGAAHL	1339	1.074.01	1.0	12
DWKPFGGSC	21	1.0506	1.0	14
ELITSCSSNV	2814	1.0506	1.0	100
EPRKQKALGL	1731	1.0491	1.0	12
EVNTSTWNLV	1659	1.0509	1.0	12
GLSAFSLHSY	2921	1.0491	1.0	11
GLSTLPGRPA	1782	1.0498	1.0	14
GLTHIDAHFL	1569	1.0498	1.0	13
GPSEGAVQWM	1912	15.0240	1.0	86
GONGGVNL	28	1.0504	1.0	13
GIVGTVVYHGA	1091	1.0504	1.0	79
GIVVCEKHL	2619	1.0504	1.0	100

## HCV-B62 Super Motif

Sequence	Position	Peptide No.	No. of Amino Acids	Sequence Frequency	Conservancy (%)
HONIVDQYI	898		10	11	79
LAGYAGAGVA	1856		10	11	79
ILGGWVAQQL	1818		10	12	86
IMAKNEVFCV	2551		10	11	79
IOYLAGLSTL	1777		10	14	100
INFQDGVVRN	2613		10	11	79
KPTLHGPTPL	1630		10	11	79
KVDTLICGFF	121		10	12	86
KVVLVNPSSA	1255		10	14	100
LFNLGGWV	1812		10	12	86
LPAILSPGIA	1887		10	13	93
LMGYIPLVGA	133		10	11	79
LPAILSPGAL	1888		10	13	93
LPGCSFSIPL	169		10	13	93
LPRGRPRGLV	37		10	13	93
LPVCDQHLEF	1553		10	12	86
LYAQATVCA	1591		10	12	86
LYDLAGYGA	1843		10	11	79
LYGGVLAALA	1667		10	12	86
LVIGVCAAI	1897		10	11	79
MLIDPSHITA	2179		10	14	100
NPGCFSFIF	168		10	13	93
NPSVAAATLGF	1260		10	14	100
PITYSTYKGF	1285		10	11	79
PLGGAARALA	143		10	11	79
POPEYDELI	2807		10	11	79
PVCOOH-LEFV	1554		10	12	86
PVNSWLMNNI	2857		10	14	100
PVYCFTSPIV	508		10	13	93
QLPCEPFDIV	2164		10	12	86
QPEKGGRKPA	2601		10	11	79
RHGUSSAFL	2918		10	11	79
PLVFFPDGLV	2611		10	11	79
RMAYWDMMMW	317		10	12	86
RMLEGQVNA	156		10	13	93
SHEYSPGEI	2826		10	11	79
SLIGRDKNOV	1051		10	12	86
SPGALVIGIV	1883		10	11	79
SQLSAPSLSKA	2268		10	11	79
SOPRRRQP	56		10	13	93
SYAATLGFEA	1262		10	14	100
TLHGPTPLLY	1622		10	11	79
TLFLNLLGGWV	1811		10	12	86
TLPALSTGGL	695		10	11	79
TLJCGFADLM	125		10	12	86

## HCV B62 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
TPCTCGSSDL	1126	10	11	79
TRLYFLGAV	1627	10	13	93
TPVNSWLGN	2856	10	12	86
TVDFSLDFIF	1466	10	12	86
VIOTLTCGFA	122	10	12	86
VLAALAAAYCL	1671	14	12	86
VLDQKETAGA	1337	10	12	86
VNPNSVAAITL	1258	10	14	100
VLTTSGGNLT	2737	10	11	79
VLVGGVILAN	1666	10	12	86
VVLNIPSVA	1256	10	14	100
VMGSSYIGFQY	2639	10	11	79
VPESDAAAFRV	1940	10	12	86
VOWMNRILAF	1819	10	14	100
WVGWVCAAIL	1699	10	10	79
WVLYGGVILAA	1665	10	12	86
YLGESSGGPL	1165	10	12	86
YLPHRPHPL	35	10	13	93
YLVTBHDYI	1136	10	11	79
YVGQDGSYF	276	10	12	66
ALVIVWYCAI	1896	11	11	79
APFGSGKSTKV	1235	11	13	93
APTLWARMILM	2869	11	11	79
ADAPPSSWQDM	1602	11	12	86
AVCTHGVAKAV	1168	11	11	79
AVOWMNRILAF	1917	11	14	100
DILAGYAGAVIA	1855	11	11	79
DLEVTSTWNL	1657	11	12	86
DLGWRVCEQMA	2617	11	13	93
DLMGTFPLVGA	132	11	11	79
DLYVTRBHDY	1134	11	12	86
DOAEVAGAFRLV	1339	11	12	86
DWKFPGGQV	21	11	12	86
EOKFKKALELL	1731	11	14	100
FISGIVNAGL	1773	11	11	79
FLAGGESESGA	1304	11	14	100
FGGGGNGGN	24	11	11	79
FOYSGSREVF	2646	11	11	100
GIVYLAGLSTL	1778	11	14	100
GLPVOOCHLIF	1552	11	12	86
GLSTLPGNPAI	1782	11	11	79
GPTPLYRIGA	1625	11	13	93
GPVYCFTPSPV	607	11	13	93
GVLAALAAVCL	1670	12	12	86
GVRVCEMAYL	2118	14	100	

## HCV I62 Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
GIVMLEDGVNY	154	12	86	
H1HNNIDVQY	696	11	79	
HWNINFSQIY	1769	13	93	
HNNIDVQY	698	11	79	
HAGPREGAVW	1910	11	79	
ILGGEWVAAQLA	1615	12	86	
ILGIGTVDDQ	1331	12	86	
LSFGSALWVGIV	1691	13	93	
KPARIIVFPL	2608	11	79	
KPTIHQGPTPL	1620	11	79	
KOKALGILLOTA	1734	12	86	
KVIDILTGCF	121	12	86	
KVLYLNPSVAA	1255	14	100	
LIAFASRGNHV	1924	14	100	
LITSCSSNVSV	2815	14	100	
LIVPPDGLGV	2612	11	79	
LLFLLADARV	726	13	93	
LLFRILGGWAA	1612	12	86	
LLPALSFGAL	1687	13	93	
LLPFRGQFLGV	36	13	93	
LLSPRGSRPSW	87	11	79	
LLWPROEMGSV	2240	12	86	
LPAILSFGALV	1888	12	86	
LPALSTGLHL	667	12	86	
LPGECSFISFL	169	13	93	
LPVCOOHLEFW	1553	12	86	
LVGGVIAAIAA	1667	12	86	
LYLNPSVAAATL	1257	14	100	
LYTAHADWIPV	1137	11	79	
LVYGWVCAAIL	1887	11	79	
NILGGWVAAQ	1815	12	86	
NITRVESENKV	2249	12	86	
NLLPAILSFGA	1866	13	93	
NPECSEFQL	168	13	93	
PITYSTYGRFL	1295	11	79	
PLEGERCPDPL	2403	13	93	
PMGFSYDTRCF	2667...	11	79	
PPFSNDOMVYKQL	1696	11	78	
PVNSWLGNIM	2857	12	86	
PVCFITPSIPW	508	13	93	
PRMVGGGVBHL	635	13	93	
RCENMGSNTRV	2244	12	88	
RVCEKMALYDV	2B21	12	86	
SIFLLAISCL	175	12	86	
SIMLTDPSHTA	2178	14	100	

## HCV\_B62\_Super Motif

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)
SPIHYPPESDA	1935	11	12	86
SCLPCEPERDV	2163	11	12	86
SVATLQFGAY	1262	11	14	100
TLGFGATMSKA	1266	11	12	86
TILENLSGAVV	1811	11	12	86
TPCTTGESSDLY	1126	12	11	79
TPGLFPVCDHL	1550	11	13	83
TPVNSWLGNII	2856	11	12	86
TVLDQAEETAGA	1336	11	12	86
VCCECYDAGCA	1521	11	11	79
VLVDILAGYGA	1852	11	11	79
VLVGSQMLALA	1666	11	12	86
VPDKGGKPKPA	2600	11	11	79
YQWMNRHIAFA	1918	11	14	100
YVCAALRIRHV	1901	11	11	79
WWLVGGVLAAL	1665	11	12	86
YLGKSSGGPLL	1165	11	12	86
YLVAYQATVCA	1590	11	12	86
YQATVCARAQIA	1594	11	11	79
YVGDQCESEVRL	276	12	86	86
YYPESDAARAV	1939	11	12	

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Table XV HCV  $\alpha$ 01 Motif with Binding Information

Sequence	Position		No. of Amino Acids	Frequency	Conservancy (%)	A <sup>+</sup> 0101
ASF CGSPY	166	26.0026	9	20	100	
DN SVL SRKY	737	20.0255	10	18	90	0.0001
FA APFT QCGY	631	20.0254	10	19	95	0.0680
GF APFT QCGY	630		11	19	95	
GRE TLEY	140		8	15	75	
GY SLNF MGY	579	2.0058	9	17	85	
HT LWKAG ILY	149	1069.04	10	20	100	0.1100
KO AFT SPY	653	20.0256	10	19	95	0.0001
LL DTAS ALY	30	1069.01	9	17	85	12.0000
LS LDV S A M F Y	415	1090.07	10	19	95	0.0150
LT FGRE TLEY	137		11	15	75	
MM WYW G P SLY	360	1039.01	10	17	85	0.0810
MS TTD L E A Y	103	2.0126	9	15	75	0.8500
NS V M L SR KY	738	2.0123	9	18	90	0.0005
P L O G K I K P Y	124	1147.12	9	20	100	
P L O G K I K P Y	124	1069.03	10	20	100	0.1700
PT TG R G T S LY	797	1090.09	9	17	85	0.2100
S A S F CG S P Y	165		9	20	100	
S L D V S A M F Y	416	1069.02	9	19	95	
S T T D L E A Y	104		8	15	75	
T G R G T S LY	798	26.0030	8	17	85	
W L S LD V S A M F Y	414	26.0051	11	19	95	
W M W Y W G P S	359	1039.06	11	17	85	0.3200
Y P A M P L Y	640	19.0014	8	19	95	
Y S L N F M G Y	580	26.0032	8	17	85	

Table XVI HCV A03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*0301
AACNVTIGER	647	10	12	86	0.0003
AARALAHGVR	147	10	11	79	
AATLGFGA	1264	0	14	100	
AATLGFGAY	1264	0	9	14	100
AAVCTRGVVA	1187	9	11	79	
AAVCTRGVAK	1187	10	11	79	
AAVCTRGVAKA	1187	11	11	79	
ACRMVTRGER	648	9	12	86	
ADGCGSGGA	1306	9	11	79	
ADGGCGSGAY	1306	10	11	79	
ADVIPVRR	1142	0	12	06	
ADVIPVRR	1142	9	11	79	
AFASRGNH	1926	0	14	100	
AGALVAFK	1865	0	12	06	
AGARLVLVA	1344	9	12	06	
AGARLVLVATA	1344	11	11	79	
AGLSTLPPNPA	1781	11	14	100	
AGVAGALVA	1862	9	12	06	
AGVAGALVAF	1862	10	12	06	
AGVAGALVAFK	1862	11	12	06	
AGMILSPR	94	0	12	06	
AGMILSPRGSR	94	11	12	06	
AGYGAAGVA	1858	0	12	06	
AGYGAAGVAGA	1050	10	12	06	
ALGILLOTA	1737	0	12	06	
ALSTGILIH	609	0	12	06	
ALSTGILIH	609	10	12	06	
ALVGVICVA	1096	9	11	79	
ALVGVICVA	1896	10	11	79	
ASLIMAFTA	1793	0	11	79	
ASQLSAPSLSK	2208	10	11	79	
ASQLSAPSLSK	2208	11	11	79	
ASRGNHVSPTH	1928	11	12	06	
ASSASOLSA	2204	10	14	100	
ATGNLPGCSF	165	10	13	93	
ATLGFGAY	1265	0	14	100	
ATLGFGAYNSIK	1265	11	12	06	
ATRKTSER	48	0	11	79	
ATVCGARAAQ	1596	9	11	79	
AVCTRGVVA	1108	0	11	79	
AVCTRGVAK	1108	9	11	79	0.0260
AVCTRGVAKA	1108	10	11	79	
AVQWMNRLLIAF	1917	11	14	100	
CAAILRRH	1903	8	13	93	

## HCV A03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'3301
CAWYELTPA	1630	9	11	79	
CGFADLMGY	120	9	13	93	
CGNTLTCY	2742	8	11	79	
CGSSDLYLVH	1130	11	11	79	
CGYRICRA	2727	8	14	100	
CLRLKGVPPLR	2941	11	12	86	
CSFSFILLA	172	9	14	100	
CSSNVNSVA	2819	8	14	100	
CSNVSYAH	2819	9	12	86	
CTCGSSDLY	1128	9	11	79	
CTRGVAKA	1190	0	11	79	0.0001
CTRGVAKAYDF	1190	11	11	79	
CTWMNISTGF	555	9	11	79	
CTWMNISTGFK	555	11	11	79	0.7600
CYOPFKGKR	2599	9	13	79	0.0009
CYOPEKGGRK	2599	10	11	79	0.0011
CYTORVDF	1462	8	12	86	
DAHFLSOTK	1574	9	14	100	0.0003
DDLVVICESA	2771	10	11	79	
DFSLOPTF	1468	0	14	100	
DGGCGGGA	1307	8	11	79	
DGGCGGGAY	1307	9	11	79	
DLICDCEH	1316	9	12	86	
DILAGYGA	1855	8	12	86	
DILAGYGAGVA	1055	11	11	79	
DLGVRVCEK	2617	9	13	93	0.0003
DLGVRVCEKMA	2617	11	13	93	
DLMGYIPLYGA	132	11	11	79	
DLVNLILPA	1883	8	11	79	
DLVVICESA	2772	9	11	79	
DLYLYTRH	1134	8	12	86	
OLYLVTRHA	1134	9	12	86	0.0003
DTLTGCFKA	124	8	12	86	
DVIFVPRR	1143	8	11	79	
EAMTRYSA	2794	8	14	100	
ECYDAGCA	1524	8	11	79	
ECYDAGCANY	1524	10	11	79	
EDLVNLILPA	1882	9	11	79	0.0004
EGAVGQHNR	1915	9	14	100	
EIPFGIGA	1377	8	13	93	
EMGGNITR	2245	8	12	86	
ETAGARLVLAA	1342	11	12	86	
ETTMRSVPF	1207	9	12	86	0.0008
EVFOQPEK	2596	9	12	86	
FOVOPEKGGR	2598	10	11	79	

HCV  $\Delta$ 03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A <sup>03</sup> 01
FCMOPERKGGRK	2590	11	1	1	7.9
FGAYMSKA	1269	8	12	8.6	
FGAYMSKAH	1268	9	12	8.6	
FGCTWNISTGF	553	11	11	7.9	
FGYGAKDVR	2554	9	12	8.6	0.00008
FISGIOYLA	1773	9	14	100	
FLAGGCGGGAA	1304	11	11	7.9	
FLLADAR	728	8	14	100	
FSYDTRCF	2670	8	11	7.9	
FTEAMTRY	2792	10	14	100	
FTGLTHIDA	1567	9	13	9.3	
FTGLTHDAH	1567	10	13	9.3	
FTGLTHIDAHF	1567	11	13	9.3	
GAARALAH	146	8	14	100	
GAARALAHGVR	146	10	14	100	
GAGV/GALVA	1861	10	12	8.6	
GAGVAGALVAF	1861	11	12	8.6	
GAHWGVL	350	8	12	8.6	
GALVGWVCA	1895	10	11	7.9	
GALVVGVCA	1895	11	11	7.9	
GARLWVLA	1345	8	12	8.6	
GARLWVLATA	1345	10	11	7.9	
GAVWMMNR	1916	6	14	100	
GAVWMMNLIA	1916	11	14	100	
GAYMSKAH	1270	8	12	8.6	
GCAYWELTPA	1529	10	11	7.9	
GCSFSFLA	171	10	14	100	
GCYTMNSTGF	554	10	11	7.9	
GDDLWVCESA	2770	11	11	7.9	
GLOGSVF	278	8	12	8.6	
GFADLWGY	129	8	13	9.3	
GFGAYMSK	1268	8	12	8.6	
GFAYMSKA	1268	9	12	8.6	
GFAYMSKAH	1268	10	12	8.6	
GFGYSPCOR	2645	9	11	7.9	
GSYDTRCF	2669	9	11	7.9	
GGAAATALA	145	8	11	7.9	
GGAAHALAI	1308	5	11	7.9	
GGCGGGAY	26	10	14	100	
GGGONGGGVY	935	8	11	7.9	
GGHYVOMA	27	9	14	100	
GGGQGGVY	1392	9	14	100	
GGHLIFCHSK	1392	11	14	100	

ICV $\Delta$ 03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*0301
GEGKPARLIVF	2605	11	11	79	
GGVLAALAA	1669	8	12	86	
GGVLAALAA	1669	9	12	86	
GGVLAALAA	1669	10	12	86	
GGYILPFR	32	8	13	93	0.0003
GGYILPFR	32	9	13	93	
GGWVAQLA	1818	9	12	86	
GIGTVLDOA	1333	9	14	100	
GIYLPNR	3037	8	11	79	
GLPYCDDH	1552	8	13	93	
GLPYCDDH	1552	11	12	86	
GLPVODDLEF	1004	0	11	79	
GLPVSAARR	968	8	11	79	
GLRDLAYA	2921	0	11	79	0.0100
GLSAFSLHSY	2921	10	11	79	
GLSTLPGNPA	1782	10	14	100	
GLTHIDAH	1569	8	13	93	
GLTHIDAHF	1569	9	13	93	
GSGKSTKVPAA	1238	10	12	86	
GSGKSTKVPAA	1238	11	12	86	
GSSDLVLVTR	1131	10	12	86	
GSSDLVLVTRH	1131	11	12	86	
GSSYGFQY	2641	8	11	79	
GTEPINAY	2063	0	11	79	
GTVLDOAETA	1335	10	14	100	
GVAGALVA	1863	0	12	86	
GVAGALVAF	1863	9	12	86	
GVAGALVARK	1063	10	12	86	0.3900
GVAKAVDF	1193	0	11	79	
GVCWTVYH	1081	8	11	79	
GVCWTVYHGA	1081	10	11	79	
GIGYILPFR	3035	10	11	79	0.0014
GGVLAALAA	1670	8	12	86	
GGVLAALAA	1670	9	12	86	0.0046
GVRAFKTSER	45	11	11	79	
GVRCCEKMA	2619	9	14	100	
GVRCCEKMA	2619	11	14	100	
GVRLLEDGINY	154	11	12	86	
GVVCAAILR	1900	9	11	79	
GVVCAAILR	1900	10	11	79	
GVVCAAILRH	1900	11	11	79	
GVVILPFR	33	8	13	93	
GVVILPFRGPR	33	11	13	93	
HADVIPVR	1141	6	11	79	
HADVIPVR	1141	9	11	79	

## HCV Δ93 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'0301
HADVIPYARR	1141	10	1.1	79	
HAPTGSK	1234	0	1.4	100	
HAPTGSKSTK	1234	11	1.3	93	
HGLSAFSLH	2920	9	1.1	79	
HGLSAFSLHSY	2920	11	1.1	79	
HGPTPLLY	1624	0	1.1	79	
HGPTPLLYR	1624	9	1.1	79	
HDAHFLSQTK	1572	11	1.4	100	
HHAHPTGSK	1232	10	1.2	86	0.5000
HIFONIVDQY	696	11	1.1	79	
HIFCHSK	1305	0	1.4	100	
HIFCHSKK	1395	9	1.4	100	0.0250
HIFCHSKKK	1395	10	1.4	100	0.0260
HMKNIFISQY	1769	11	1.3	93	
HSKKKCDELA	1400	10	1.4	100	
HSKKKCDELAA	1400	11	1.4	100	
HSVSGEINR	2926	10	1.1	79	0.0004
HTFGCVPCVR	222	10	1.1	79	
HWGPGEKA	1910	0	1.1	79	
IAFASRGNH	1925	9	1.4	100	0.0003
IDAHFLSQTK	1573	10	1.4	100	
IDLTTCGF	123	8	1.2	86	
IDLTCGFA	123	9	1.2	86	
IFCHSKKK	1397	8	1.4	100	
IGTVLOCK	1334	0	1.4	100	
IGTVDOAETA	1334	11	1.4	100	
IMDCECH	1317	8	1.2	86	
ILAGYGAGVA	1656	10	1.1	79	
ILGGWVAA	1016	8	1.2	86	
ILGGWVAAQLA	1816	11	1.2	86	
IGIGTVDOA	1331	11	1.2	86	
IMAKNEVF	2591	8	1.2	86	
ISGIOYLA	1774	0	1.4	100	
ITRVESENK	2250	9	1.2	86	0.0150
ITSCCSANVSA	2816	11	1.4	100	
ITWGADTA	989	8	1.2	86	
ITWGADTA	989	9	1.2	86	
ITYSTYK	1296	8	1.2	86	
ITYSTYKGF	1296	9	1.2	86	
ITYSTYKFLA	1296	11	1.1	79	
IVDVQVLY	701	0	1.2	86	
IVFPDQVRA	2613	9	1.1	79	0.0036
IVGGVYLLPRA	30	10	1.3	83	0.0008
KALGLLOTA	30	11	1.3	93	
		9	1.2	86	
					1736

ILCV $\Delta$ 03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*0301
KCDELAAK	1404	8	12	86	
KGYGAKDVR	2553	10	12	86	
KGGRH1LF	1391	8	11	79	
KGGHLIFCH	1391	10	11	79	
KGGRKPAH	2604	6	11	79	
KLGVPPLR	2944	6	12	86	
KSTKVKPAA	1241	6	12	86	
KSTKVKPAA	1241	9	12	86	0.0009
KSTKVKPAA	1241	10	12	86	
KSTKVKPAA	1241	11	11	79	
KTKRANTNR	10	0	12	86	
KTKRANTNR	10	9	12	86	0.0110
KTSERSQPR	51	9	13	93	0.1600
KTSERSQPR	51	11	12	86	
KVDTLTCGF	121	10	12	86	
KVDTLTCGF	121	11	12	86	
KVILVLPNSVA	1255	10	14	100	
KVILVLPNSVA	1255	11	14	100	
KVPAAYAA	1244	8	11	79	
LAGGGCGGAA	1305	10	11	79	
LAGGGCGGAY	1305	11	11	79	
LAEORKOK	1729	6	12	86	
LAEORKOK	1729	9	12	86	
LAGYGAGVA	1057	9	11	79	
LAGYGAGVGA	1057	11	11	79	
LCFCYDGCAC	1522	10	11	79	
LDQNETGAG	1330	9	12	86	
LDQNETGAG	1330	10	12	86	
LFLLIADAR	727	8	14	100	
LFLLIADAR	727	9	14	100	
LFNLLGGWAA	1013	10	12	86	
LFNLLGGWAA	1013	11	12	86	
LFIFSPR	290	8	11	79	0.0810
LGFGAYMSK	1267	9	12	86	
LGFGAYMSKAH	1267	10	12	86	
LGGAARALA	144	9	11	79	
LGGAARALA	144	10	11	79	
LGGAARALAH	1617	10	12	86	
LGGVWAAQLA	1332	10	13	93	
LGIVATDK	44	8	12	86	
LGIVRCEK	2616	8	14	100	
LGIVRCEK	2616	10	14	100	
LIAFASRGRNH	1924	10	14	100	
LIEANLLWV	2235	9	12	86	0.0008

## HCV\_A03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A-0301
LIFCHISKK	1390	0	1.4	100	0.5400
UFOHSKKK	1396	9	1.4	100	
LINTNGSMH	414	9	1.1	79	
LNFPOLGVR	2612	10	1.1	79	0.0003
LLAPITAY	1030	8	1.4	100	
LLFLFLLADA	726	9	1.4	100	0.0016
LLFLLLADAR	726	10	1.4	100	
LLENLIGKWA	1812	11	1.2	86	0.0003
LLPAILSPGA	1687	10	1.3	93	
LLPARGPR	36	8	1.3	93	
LLSPRGSR	97	8	1.2	86	
LMGYIPLVGA	133	10	1.1	79	
LSAFESLHSY	2922	9	1.1	79	0.0002
LSAPSLSKA	2211	0	0	11	
LSNSLPH	2479	0	1.2	86	
LSNSLRAHH	2479	9	1.2	86	0.0003
LSTGULHHL	690	9	1.2	86	
LSTLPGNPA	1783	9	1.4	100	
LTCGFADLNGY	126	11	1.2	86	
LTDPSPHTA	2180	9	1.4	100	
LTHDAHF	1570	8	1.3	93	
LTSMILTOPSH	2176	10	1.3	93	
LWACATVCA	1591	10	1.2	86	
LWYDATVCAH	1591	11	1.1	79	
LVDILAGY	1053	0	1.1	79	
LVDILAGYGA	1053	10	1.1	79	
LVGGVLA	1667	0	1.2	86	
LVGGVLAALAA	1667	10	1.2	86	
LVLNPSVA	1257	8	1.4	100	
LVLNPSVAA	1257	9	1.4	100	
LVNGVWCA	1897	8	1.1	79	
LVGVVCAA	1897	9	1.1	79	
LWVICESA	2773	8	1.1	79	
MGFSYDTR	2668	8	1.1	79	
MGFSYDTRCF	2660	10	1.1	79	
MGSYYGFOY	2640	9	1.1	79	
MGYIPLVGA	134	9	1.1	79	
MILMTHFF	2876	0	1.2	66	
MLTDPSPHTA	2179	10	1.4	100	
MSTNPKPORK	1	9	1.1	79	
NOGYRRCR		10	1.1	79	
NCGYRRCR	2726	8	1.1	79	
NCSYPGH	2726	9	1.1	79	
	305	8	1.1	79	

## ICV A01 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*0301
NFISGQY	1772	0	14	100	
NFISGQYLA	1772	10	14	100	
NGCWCWVY	1080	0	11	79	
NGCWCWVYH	1080	9	11	79	
NGCWCWVYHGA	1080	11	11	79	
NILGGWVA	1815	8	12	86	
NILGGWVA	1815	9	12	86	
NTRVESENK	2249	10	12	86	0.0010
NIVDVOQLY	700	9	12	86	0.0005
NILPAILSPGA	1086	11	13	93	
NLPGCSFSIF	168	10	13	93	
NTCVTQTVDF	1460	10	12	86	
NTNARRPODK	14	10	11	79	0.0010
NTNARRPODKF	14	11	11	79	
NTPLGLPVCOOH	1549	11	13	93	
PAILSPGA	1089	8	13	93	
PAISSTGJH	608	9	12	86	
PAISSTGJH	608	11	12	86	
PGSGWMLR	1976	8	11	79	
PGTGSSSLY	1127	10	11	79	
PDGGRVCEK	2616	10	13	93	
PGALVVGVVCA	1094	11	11	79	
PGCSFSIF	170	8	14	100	
PGCSFSIFLA	170	11	14	100	
PGCVPVCV	224	0	12	66	
PGEGAVQWMAA	1913	11	13	93	
PGENRVA	2932	0	11	79	
PGERPSGMF	1509	9	12	86	
PGGCGGGVY	25	11	14	100	
PGPWCODH	1551	9	13	93	
PGYPWFLY	79	8	14	100	
PTVSTYK	1295	9	11	79	
PTVSTYK	1295	10	11	79	
PLGGAAARAH	143	8	13	93	
PLGGAAARAH	143	8	11	79	
PLLYRFLGA	1628	9	11	79	
PMGFSYDTRF	2667	11	11	79	
PMGFSYDTRF	2667	11	13	93	
PSPVWNGTDR	514	11	13	93	
PSVATLGF	1261	9	14	100	
PSVATLGF	1261	11	14	100	
PSWDDWKK	1607	8	11	79	
PTDCFRKH	507	0	13	93	
PTDFRRRSR	109	9	12	86	0.0008

## HCVΔ03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'0301
PTGSGKSTK	1236	9	13	93	0.0002
PTHYVPESDA	1936	10	12	86	
PTHYVPESDAA	1936	11	12	86	
PTLHGPTPFLY	1621	11	11	79	
PTPLLYRIGA	1626	10	13	93	
PVQOCHLEF	1554	9	12	86	
PVWVGTIDR	516	9	13	93	0.0008
QAEETGAR	1340	8	12	86	
QATVCARA	1595	8	13	93	
QATVCARAQ	1595	10	11	79	
QVGGYVLPF	29	11	13	93	
QLTESPR	209	9	12	86	
QLTFSPRR	209	9	11	79	0.7500
QLTRPOA	336	8	12	86	
QLSAPSILK	2210	8	11	79	
QLSAPSILKA	2210	9	11	79	
QYDVSLSIDPTF	1465	11	12	86	
RAAVCTRGVA	1186	10	11	79	
RAAVCTRGVAK	1186	11	11	79	
RALAHGVIR	149	8	14	100	
RATRKTSEH	47	9	11	79	
FGINHVSPTH	1930	9	12	86	0.0003
FGINHVSPTH	1930	10	12	86	0.0003
FGPFLGVIR	40	9	13	93	
FGPFLGVRA	40	9	13	93	
FGPFLGVATR	40	11	11	79	
FGTROPPIK	59	9	13	93	0.0120
FGSLLSPR	1154	8	12	86	
FGVAKAVDF	1192	9	11	79	
FGVAKAVDF	1192	9	11	79	
FGVFRATR	43	8	11	79	0.9400
FGVFRATR	43	9	11	79	
FLHGSAF	2918	8	12	86	
FLHGSAFSLH	2918	11	11	79	
FLIAFASH	1923	8	14	100	
FLIAFASRGNH	1923	11	14	100	
FLVFPFDLGVR	2611	11	11	79	
FLGVFRATR	43	9	12	86	
FLLGSAF	1029	8	12	86	2.7000
FLLGSAFSLH	1029	9	12	86	
FLLAPITAY	1029	8	12	86	
FLVFLATA	1347	8	12	86	
FLMILMTHF	2875	8	12	86	
FLMILMTHFF	2875	9	12	86	
FLMVGGVHEH	635	9	14	100	0.7200
FLMVGGVHEH	635	10	14	100	
RSQCPGGR	55	8	13	93	
RVCEKMMALY	9	14	100	0.1800	

HCV A03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'0301
RVLEDGVNYA	156	1174-17	9	12	86
SAFLSHY	156		10	12	86
SASQSLAPSILK	2923		8	11	79
SCSSNNVSAH	2207		11	11	79
SDLYLVTR	2018		9	14	100
SDLYLVTRH	2018		10	12	86
SDLYLVTRH	1133		8	12	86
SDLYLVTRH	1133		9	12	86
SFSIFLLA	1133		10	12	86
SGKSTKVPAA	173		6	14	100
SGKSTKVPAA	1239		9	12	86
SGKSTKVPAA	1239		10	12	86
SMLTQPSH	1239		11	12	86
SMLTQPSHITA	2178		6	14	100
SSASQSLSA	2206		11	14	100
SSDLYLVRH	2206		0	14	100
SSDLYLVRH	1132		9	12	86
SSDLYLVRH	1132		10	12	86
SSNVSVAH	2820		6	12	86
SSSAQSLSA	2205		9	14	100
STGLIHLH	691		8	12	86
STKVPAAAY	1242		0	12	86
STKVPAAAY	1242		0	12	86
STKVPAAAYA	1242		9	12	86
STLPGNPA	1704		10	11	79
STNPKPQR	2		0	14	100
STNPKPQRK	2		0	11	79
STNPKPQRKT	2		9	11	79
STWVLGGVLA	1663		11	11	79
STYGKELA	1298		8	12	86
SVAAATLGFL	1262		8	14	100
SVAAATLGFGA	1262		10	14	100
SVAAATLGFGAY	1262		11	14	100
TGARLWLA	1343		10	12	86
TGFFADLMGY	127		10	13	93
TGSSDOLY	1129		6	11	79
TCVTOVDF	1461		9	12	86
TDPPRPSR	1110		0	12	86
TDPSHTA	2101		0	14	100
TGEIPFYGK	1375		9	11	79
TGEIPFYGKA	1375		10	11	79
TGLTHIDA	1568		6	13	93
TGLTHIDAH	1568		9	13	93
TGLTHIDAH-F	1568		10	13	93

ICV  $\Delta$ 03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A <sup>03</sup> 01
TGNLPGCSF	166	9	1.3	9.3	
TGSGKSTK	1237	6	1.3	9.3	
TGSGKSTKVPAA		11	1.2	8.6	
TIMAKNEVF	2550	9	1.1	7.9	
TLGFAYMSK	1268	10	1.2	8.6	0.0810
TLGFAYMSKA	1266	11	1.2	8.6	
TLGKPTPLLY	1622	10	1.1	7.9	0.0890
TLGKPTPLYR	1622	11	1.1	7.9	
TLPLSTGLIH	686	11	1.1	7.9	
TLWARMILMTH	2871	10	1.4	100	
TSSSNVSYAH	2017	11	1.2	8.6	
TSSSNVSYAH	2017	11	1.2	8.6	
TSERSOPR	52	8	1.3	9.3	0.0003
TSERSOPRGR	52	10	1.2	8.6	
TSERSOPRGRG	52	11	1.2	8.6	
TSLTGRDK	1050	8	1.2	8.6	
TSMLTDPSH	2177	9	1.3	9.3	0.0003
TTIMAKNEVF	2589	10	1.1	7.9	
TMRSVPF	1208	8	1.2	8.6	
TVCARAQA	1597	8	1.1	7.9	
TVDFSLDPTF	1466	10	1.2	8.6	
TVLDQETA	1336	9	1.4	100	
TVLDQETAAGA	1336	11	1.2	8.6	
VANLGFQA	1203	9	1.4	100	
VANLGFQAY	1203	10	1.4	100	
VAGALVAF	1064	8	1.2	8.6	0.2400
VAGALVAFK	1064	9	1.2	8.6	
VAYQATVCA	1592	9	1.2	8.6	
VAYQATVCA	1592	10	1.1	7.9	0.0005
VAYQATVCA	1592	11	1.1	7.9	
VCAAILRR	1902	6	1.1	7.9	
VCAAILRRH	1902	9	1.1	7.9	
VCEKMLY	2622	8	1.4	100	
VOGRVYCF	505	8	1.3	9.3	
VOQATVCA	1592	11	1.2	8.6	
VCAAILRR	1902	6	1.1	7.9	
VCTRGVAKA	1109	9	1.1	7.9	
VGMVYHGA	1082	9	1.1	7.9	
VDLSDLPTF	1467	9	1.4	100	
VDLAGYGA	1555	8	1.1	7.9	
VCTRGVAK	1109	9	1.1	7.9	
VCTRGVAKA	1109	9	1.1	7.9	
VGMVYHGA	1082	9	1.1	7.9	
VDFSLDPTF	1467	9	1.4	100	
VDFSLDPTF	1467	10	1.3	9.3	
VDFYPLWHY	614	6	1.2	8.6	
VFCVQPEK	2597	11	1.1	7.9	
VFCVQPEKGR	2597	11	1.1	7.9	
VFPDLGVR	2614	0	1.1	7.9	

IICV $\Delta$ 03 Motif with Binding Information

Position	Sequence	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A $\cdot$ 0301
1566	VFTGLTHIDA	10	1.3	9.3	
1566	VFTGLTHIDAH	11	1.3	9.3	
277	VGDQCSMVF	9	1.2	8.6	
1668	VGGVLAALA	9	1.2	8.6	
1668	VGGVLAALAA	10	1.2	8.6	
1668	VGGVLAALAY	11	1.2	8.6	
31	VGGVYLPR	9	1.3	9.3	0.0003
31	VGGVYLPRR	10	1.3	9.3	0.0007
3036	VGYLPLNR	9	1.1	7.9	
1009	VGVVCAILR	10	1.1	7.9	
1099	VGVVCAILRR	11	1.1	7.9	
122	VIDLTCGF	9	1.2	8.6	
122	VIDLTCGFA	10	1.2	8.6	
1671	VLAALAY	0	1.2	8.6	
1521	VLCECYDA	8	1.3	9.3	
1521	VLCECYDAGCA	11	1.1	7.9	
1337	VLDQAEATA	8	1.4	100	
1337	VLDQAEATAGA	10	1.2	8.6	
1337	VLDQAEATAGR	11	1.2	8.6	
157	VLEDGVRY	8	1.2	8.6	
157	VLEDGVNYA	9	1.2	8.6	
1250	VLNPSVVA	0	1.4	100	
2175	VLTSMLTDPSH	11	1.3	9.3	
1052	VLVDILAGY	9	1.1	7.9	
1052	VLVDILAGYA	11	1.1	7.9	
1666	VLVGGVLA	0	1.2	8.6	
1666	VLVGGVLAALAA	9	1.2	8.6	
1666	VLVGGVLAALAY	11	1.2	8.6	
1256	VLVLPNSVA	9	1.4	100	
1256	VLVLPNSVAA	10	1.4	100	
2639	VMGSSYGF	0	1.1	7.9	
2639	VMGSSYGFQY	10	1.1	7.9	
1138	VTRHADWIPVR	11	1.1	7.9	
1901	VVCAAIR	8	1.1	7.9	
1901	VVCAAIRR	9	1.1	7.9	
1901	VVCAAIRRRH	10	1.1	7.9	
1098	VVGVYCAA	0	1.1	7.9	
1098	VVGVYCAILR	11	1.1	7.9	
517	VNGITDR	0	1.3	9.3	
93	WAGWLLSPR	9	1.2	8.6	
1766	WAKHAWNF	8	1.2	8.6	
76	WAQPGYWPWLY	11	1.2	8.6	
2873	WARMILMTH	9	1.2	8.6	
2873	WARMILMTHF	10	1.2	8.6	
	WARMILMTHFF	12	1.1	8.6	

## HCV\_A03 Motif with Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'0301
WGPTDPARR	107	0	1.2	0.6	
WGPTDPARR	107	9	1.2	0.6	
WGPTDPARRS	107	11	1.2	0.6	
WLSPRGSR	96	9	1.2	0.6	0.0008
WMNRLLAF	1920	8	1.4	100	
WMNRLLAF	1920	9	1.4	100	0.0003
WMNRLLAFASR	1920	11	1.4	100	
WMNRLLAFASR	557	9	1.1	79	0.0530
WMNSTGFTK	1665	9	1.2	86	
WWLVGGVLA	1665	10	1.2	86	
WWLVGGVLA	164	11	1.2	86	
YATGNLPGCSF	1526	0	1	79	
YDAGCCAWY	1315	10	1.2	86	
YDIIICDECH	1860	6	1.2	86	
YGAGVAGAG	1860	11	1.2	0.6	
YGAGVAGALVA	2644	10	1.1	79	
YGFQYSFGCR	35	9	1.3	83	0.0054
YLPRRGPR	1590	11	1.2	86	
YLVAYAOATVCA	2930	6	1.1	79	
YSPGEINR	2930	10	1.1	79	
YSPGEINRVA	2648	9	1.1	79	
YSPGCRMEF	1298	9	1.2	86	
YSTYIGKFLA	276	10	1.2	100	
YVGDLCGSVF	637	8	1.4	100	
YVGGVBR	1939	8	1.2	86	
YVPESDAA	1939	9	1.2	86	
YVPESDAAA	1939	10	1.2	0.6	0.0003
YVPESDAAAR	567	3			

Table XVII IICV All Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'1101
AACNWTGER	647	10	12	86	0.0140
AARIALAHGVR	147	10	11	79	
AATLGFGRAY	1264	9	14	100	
AAVCTRGVAK	1107	10	11	79	
ACNWTRGER	648	9	12	86	
ADGCCGGAY	1306	10	11	79	
ADVIPVVR	1142	6	12	86	
ADVIPVRR	1142	9	11	79	
AFASRGNH	1926	6	14	100	
AGALVAFK	1085	8	12	86	
AGVAGALVFK	1062	11	12	86	
AGWLLSPF	94	8	12	86	
AGWLLSPGRSR	94	11	12	86	
ALSTGILH	689	8	12	86	
ALSTGILHILH	609	10	12	86	0.0027
ASQSLAPSILK	2209	10	11	79	
ASRGNHVSPTH	1928	11	12	86	
ATLGFGAY	1265	9	14	100	
ATLGFGAYMSK	1265	11	12	86	
ATRKTSER	48	8	11	79	
AVCTRGVAK	1169	9	11	79	0.0250
CAAILRRH	1903	8	13	93	
CGFADQMGY	128	9	13	93	
CGNDLTCY	2712	9	11	79	
CGSSDLVLTIR	1130	11	11	79	
CLRKLGVPPLR	2941	11	12	86	
CNCSYPGH	304	9	11	79	
CNIVTRGER	649	8	12	86	
CSSNVSYAH	2819	9	12	86	
CTCGSSDLY	1128	9	11	79	0.0063
CTTMNINSTGFTK	565	11	11	79	0.7500
CVPQPKGGK	2599	9	11	79	0.0005
DAHFLSOTK	1574	10	11	79	0.0008
DGCGSGRAY	1307	9	10	100	0.0005
DILICDECH	1316	9	12	86	
DLGWVRCERK	2617	9	13	93	0.0002
DLYLYTHH	1134	8	12	86	
DMPVARR	1143	8	11	79	
ECDYOGCAWY	1524	10	11	79	0.0014
EGANQNMNR	1915	9	14	100	
EMGGNITR	2245	8	12	86	
EVFCVQPER	2596	9	12	86	0.0270
FWQPKGGK	2598	10	11	79	
FWQPKGGK	2598	11	11	78	

## LICV A11 Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'1101
FGAYMSKAH	1269	9	1.2	86	
FGYGAQDVR	2554	9	1.2	86	0.0005
ELLADAR	728	8	1.4	100	
FTTEAMTRY	2792	8	1.4	100	
FTGLTHIDAH	1567	10	1.3	93	
GAARALAH	146	8	1.1	79	
GAARALAHGVR	146	11	1.1	79	
GAVCWAHNR	1916	8	1.4	100	
GAYMSKAH	1270	8	1.2	86	
GFADLMGY	129	0	1.3	93	
GFGAYMSK	1260	6	1.2	86	
GFGAYMSKAH	1260	10	1.2	86	
GIGVSPQR	2645	9	1.1	79	
GGAAARALAH	145	9	1.1	79	
GCGSGGAY	1308	0	1.1	79	
GGGGCGGAY	26	10	1.4	100	
GGQNGGYY	27	9	1.4	100	
GGRHLFLPSK	1392	9	1.4	100	
GGVLAALAY	1669	11	1.4	100	
GGYILPRA	32	10	1.2	86	
GGYLLPRA	32	8	1.3	93	
GILLLPNA	3037	0	1.1	79	
GLPVOODH	1552	0	1.3	83	
GLPVSSARR	1004	0	1.1	79	
GLSAFSLH	2921	0	1.1	79	
GLSAFSLHSY	2021	10	1.1	79	0.0005
GLTHIDAH	1569	0	1.3	93	
GNFNSPHT	1931	0	1.2	86	
GNHVSPTHY	1931	9	1.2	86	
GNITRVESENK	2248	11	1.2	86	
GSSDLVLVTR	1131	10	1.2	86	
GSSDLVLVTRH	1131	11	1.2	86	
GSSYGFQY	2641	8	1.1	79	
GTIPINAY	2063	8	1.1	79	1.4000
GVAGALVARK	1863	10	1.2	86	
GVCGWVYH	1081	8	1.1	79	0.0140
GVGKLLPNA	3035	10	1.1	79	0.0110
GVLAALAY	1670	9	1.2	86	
GVATRKTSER	45	11	1.1	79	
GVRCVCKMAY	2619	11	1.4	100	
GVWLEDGANN	154	11	1.2	86	
GVVCAAILR	1900	9	1.1	79	
GVVCAAILRH	1900	10	1.1	79	

## ICV All Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*1101
GYLLPAR	33	8	1.3	9.3	
GYLPERGPR	33	11	1.3	9.3	
HADVIPVR	1141	9	1.1	7.9	
HADVIPVRA	1141	10	1.1	7.9	
HADVIPVRR	1141	8	1.4	100	
HAPTGSK	1234	11	1.3	9.3	
HAPTGSKSTK	1234	11	1.1	7.9	
HGLSAFSLH	2920	9	1.1	7.9	
HGLSAFSLHSY	2920	11	1.1	7.9	
HGPTPLLY	1624	8	1.1	7.9	
HGPTPLYN	1624	9	1.1	7.9	
HDAHFSLQTK	1572	11	1.4	100	
HLHAPTGSK	1232	10	1.2	8.6	0.0024
HJLQHNDVQY	696	14	1.1	7.9	
HJFCHSK	1395	0	1.4	100	
HJFCHSKK	1395	9	1.4	100	0.0006
HJFCHSKKK	1395	10	1.4	100	0.0002
HMWNFISGQY	1769	11	1.3	9.3	
HSYSPGEINR	2920	10	1.1	7.9	
HTPGCVPFCVA	222	10	1.1	7.9	
IAFASRGNH	1925	9	1.4	100	0.0012
IAHFSLQTK	573	10	1.4	100	0.0003
IFCHSKKK	1397	8	1.4	100	
IIICDECH	1317	8	1.2	8.6	
INTNGSMW	415	8	1.1	7.9	
ITRVESENK	2250	9	1.2	8.6	
ITYSTYKG	1296	8	1.2	8.6	
IVDQYLY	701	0	1.2	8.6	
IVFPDLGAVR	2613	9	1.1	7.9	0.0044
IVGGVYLLPFA	30	10	1.3	9.3	0.0056
IVGGVYLLPFR	30	11	1.3	9.3	
KCDELAAK	1404	8	1.1	7.9	
KFGYGAKDVR	2553	10	1.2	8.6	
KGGPHLIFCH	1391	10	1.1	7.9	
KGGRKPAR	2604	8	1.1	7.9	
KLGVPPRL	2944	8	1.2	8.6	
KNEFQVQPEK	2594	11	1.1	7.9	
KSTKVPAAV	1241	9	1.2	8.6	0.0001
KTKANTNR	10	8	1.2	8.6	
KTKRNTNAR	10	9	1.2	8.6	0.0100
KTSERSOPR	51	9	1.3	9.3	0.0640
KTSERSQPRQR	51	11	1.2	8.6	
LADGGCGGGAY	1306	11	1.1	7.9	
LAEOFRK	1729	8	1.2	8.6	
LDQAEATGAR	1338	10	1.2	8.6	

## HCV 3' UTR Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'1101
LFLLLADAR	727	9	14	100	
LFTFSPRR	290	6	11	79	
LGFGAYMSK	1267	9	12	86	0.2900
LGGAARALAH	144	11	12	86	
LGVRATHK	44	10	11	79	
LGVRVCEK	2618	8	12	86	
LIAFASRGNH	1924	8	14	100	
LIEANLLWR	2235	10	14	100	
LIFCHSKIK	1396	9	12	86	0.0005
LIFCHSKIK	1390	8	14	100	0.1900
LINTNGSMW	414	9	14	100	
LWPFFDLGIVR	2612	10	9	79	0.0001
LLAPITAY	1030	8	14	100	
LLFLLLADAR	726	10	14	100	
LPPRGPR	36	0	13	93	
LSSPRGSR	97	8	12	86	0.0002
LSAFSLHSY	2922	9	11	79	
LSNSLRLHH	2479	8	12	86	
LSNSLRLHH	2479	9	12	86	0.0001
LSTGGLHLH	690	9	12	86	
LTGCFADLGMV	126	11	12	86	
LTSMLTDPSH	2176	10	13	93	
LVAYQAVCAH	1591	11	11	79	
LVDILAGY	1053	0	11	79	
MGFSDYDTR	2660	0	11	79	
MGSYYGFOY	2640	0	11	79	
MNRLIAFASR	1921	9	11	79	
MNSTGFTK	650	10	14	100	
MSTNPKPQR	1	0	11	79	
MSTNPKPQRK	1	9	11	79	
NCGGYFRCR	2726	8	11	79	
NCISYPGH	305	8	11	79	
NPISGICY	1772	0	14	100	
NGVCMVY	1080	8	11	79	
NGVCMVYH	1080	9	11	79	
NTRVESENK	2249	10	12	86	0.0062
NIVDVOYLY	700	9	12	86	0.0140
NTNRPAPDVK	14	10	11	79	0.0007
NTPGGLVQODH	1549	11	13	93	
PALSTGLIH	688	9	12	86	
PALSTGLHLH	688	11	12	86	
PGSGSMR	1976	8	11	79	
PCTGASSDLY	1127	10	11	79	
PDLGVVCEK	2616	10	13	93	

## ICV All Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A 1101
PGCVPCVRA	224	0	12	06	
PGEGAVGMNN	1913	11	13	93	
PGEQAVGGMYY	25	11	14	100	
PGLPVQODH	1551	9	13	93	
PGYPMWFLY	79	8	14	100	
PIYSTYSGK	1295	9	11	79	
PLGGAARALAH	143	11	11	79	
PMGFSYDTR	2667	9	11	79	
PNIRIGVRA	1281	8	13	93	
PSPVWNGTIDR	514	11	13	93	
PSNDQWMK	1607	0	11	79	
PTDCFRKH	507	0	13	93	
PTDPARASH	109	9	12	06	0.0005
PTGSGKSTK	1236	9	13	93	0.0001
PTLHGPTPLLY	1621	11	11	79	
PVNGVTTDR	518	9	13	93	0.0005
QAEETAGAAR	1340	8	12	86	
QIVGGYLLPRA	29	11	13	93	
QLFTESPR	289	0	12	86	
QLFTFSPRR	289	9	11	79	0.0330
QLSAPSILK	2210	8	11	79	
QNIIVDVOY	699	0	11	79	
QNIIVDVOYLY	699	10	11	79	
RAAVVCTGRVAK	1100	11	11	79	
RAJAHGAVR	119	0	14	100	
RATRKTSEER	47	9	11	79	
RGFMNVSPTH	1930	8	12	86	0.0001
RGHN4VSPTH	1930	10	12	86	0.0001
RGPRLGAVR	40	8	13	93	
RGPRLGVRAIR	40	11	11	79	
RGRROPKPK	59	9	13	93	0.0017
RGSLLSPR	1154	8	12	86	
RIGVYBATH	43	8	11	79	0.0290
RIGVYRATRK	43	9	11	79	
RIGHGLSAFSLH	2918	11	11	79	
RILJAFASR	1923	8	14	100	
RILJAFASRGNH	1923	11	14	100	
RILVFRPLGVR	2611	11	11	79	
RILLAPITAY	1029	9	12	86	0.0270
RIMVGGVERH	635	9	14	100	0.0200
RIMVGGVERH	635	10	14	100	
RINTRRPPDVK	13	11	11	79	
RSCPFRGPR	55	8	13	93	0.5000
RVCERMALY	2421	9	14	100	0.0068
RVLEDGVNN	156	12	86		

## ICV All Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A'1101
SAFSLHSY	2923	8	11	79	
SASQSLAPSFK	2207	11	11	79	
SCSSNVSAH	2818	10	12	86	
SDLYLVTR	1133	9	12	86	
SDLYLVTRAH	1133	9	12	86	
SGKSTKVPAAY	1239	11	12	86	
SMLTDPSH	2178	9	14	100	
SNSLAAHH	2400	8	12	86	
SSDLYLVTR	1132	9	12	86	0.0044
SSDLYLVTRAH	1132	10	12	86	0.0013
SSNVSVAH	2020	8	12	86	
STGLIHLH	691	8	12	86	
STKVKAAV	1242	8	12	86	
STNPKPQR	2	8	11	79	
STNPKPQRK	2	9	11	79	
SV/AATLGFGAY	1262	11	11	79	
TCGFADLMGY	127	10	14	100	
TCGSSDLY	1129	6	11	79	
TDPHRSR	110	8	12	86	
TGEIPFGK	1375	9	11	79	
TGLTHIDAH	1568	9	13	93	0.0001
TGSGKSTK	1237	0	13	93	
TLGEGAYMSK	1266	10	12	86	0.0010
TLHGPTPLLY	1622	10	11	79	0.0007
TLHGPTPLLYA	1622	11	11	79	
TLPALSTGQH	680	11	11	79	
TLWARMILMTH	2871	11	11	79	
TNPPIPK	3	8	11	79	
TNPKPORKTK	3	10	11	79	
TNPKPORKTKR	3	11	11	79	
TNPQDQK	15	9	11	79	
TSCESSNVSAH	2817	11	12	86	
TSERSOPR	52	6	13	93	0.0001
TSERSOPRKT	52	10	12	86	
TSERSOPRGR	52	11	12	86	
TSLTGDK	1050	6	12	86	
TSMLTDPSH	2177	9	13	93	100
VAATLGFGAY	1263	10	14	100	0.8900
VAGALVAFK	864	9	12	86	0.8900
VAYQATVCAQ	1592	10	11	79	0.0038
VCAAILRR	1802	8	11	79	
VCAAILRRH	1902	9	11	79	
VCEKMLAY	2622	8	14	100	
VCTRGVAK	1169	0	11	79	

## ICV All Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*1101
VDYPYFLWH	614	9	13	93	
VDYPYFLWHY	614	10	13	93	
VFCVQPEK	2597	8	12	86	
VFOQPERGGR	2614	11	11	79	
VFPDLGVR	1566	8	11	79	
VFTGLTHIDAH	1566	11	13	93	
VGGVLAALAY	1668	11	12	86	
VGGVYLLPR	31	9	13	93	0.0019
VGGVYLLPR	31	10	13	93	
VGYLILPNA	3036	9	11	79	0.0100
VGVYCAILRH	1099	10	11	79	
VGVVCAAILRA	1099	11	11	79	
VLAALAAV	1671	8	12	86	
VLDQAEFAGAR	1337	11	12	86	
VLEDGVNY	157	6	12	86	
VLTSMLTDPSH	2175	11	13	93	
VLVDILAGY	1052	9	11	79	
VMESSYGFQY	2639	10	11	79	
VTRHADVIPVR	1138	11	11	79	
VVCAAILR	1901	6	11	79	
VVCAAILTR	1901	9	11	79	
VVCAILRHH	1901	10	11	79	
VVGVCAAILR	1099	11	11	79	
VVGTTOR	517	6	13	93	
WAGWLLSPR	93	9	12	86	
WAGQGPYPMWL	76	11	12	86	
WARMILMTI	2073	9	12	86	
WGFTOPTRIA	107	0	12	86	
WGFTDPKPRR	107	9	12	86	
WGFTOPRSH	107	11	12	86	
WLSPRGSR	96	9	12	86	
WMNRLIAFASR	1920	11	14	100	0.0810
WMNSTGFTK	557	9	11	79	
WNFISGQY	1771	9	14	100	
YDAGCAYW	1526	6	11	79	
YDIICDECH	1315	10	12	86	
YGFQYSPGQR	2644	10	11	79	
YLLPARGPH	35	9	13	93	0.0005
YSRGENR	2930	8	11	79	
YVGCVBHR	637	0	14	100	
YVPESDAAAR	1939	10	12	86	0.0001

Table XVIII HCV A24 Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*2401
AWDMMNNW	319	8	12	0.6	
AYAAQGQYKVL	1248	10	11	79	
AYYRGGLDVSIV	1421	11	14	100	
CYDAGCAW	1525	8	11	79	
CYDAGCAWEL	1525	11	11	79	
DESLDPFIF	1468	8	14	100	
DESLDPITFI	1468	10	14	100	
FWAKHMMWNF	1765	9	12	86	6.9000
FWAKHMMWNF1	1765	10	12	86	
GFADLMGNI	129	9	13	93	
GFADLMGQPL	129	11	11	79	
GESYDGFCC	2669	9	11	79	
GWRLLAPI	1027	8	11	79	
GYGAGVAGAL	1059	10	12	86	0.0003
GYPLVGPPL	135	10	11	79	0.0057
GYRRCRASSVL	2720	11	12	86	
HMWNFISGIL	1769	9	13	93	
IFLLALLSCL	176	10	12	86	
IMANNEVE	2591	8	12	86	
KFPGCGQI	23	8	13	93	
LFNLGGGW	1813	8	12	86	
LWARMILMTHF	2872	11	12	86	
LWRCRERGQN	2241	10	12	86	
LYLVTRHADVI	1135	11	11	79	
MWNFISGI	1770	8	14	100	
MWNFISGQYL	1770	11	14	100	
MYVGGVEHHL	6336	10	13	93	0.0270
NFISGQYCL	1772	9	14	100	0.0170
PMGFSYDTCF	2667	11	11	79	
QFKOKALGL	1732	9	12	86	
QFKOKALGIL	1732	10	12	86	
QWMNRLIWF	1919	9	14	100	
QYLAGLSTL	1778	9	14	100	
QYSPGQFVEF	2647	10	11	79	0.0180
CYSQPCQFVEF	3117	10	11	79	
FIMAWDMIMNNW	2075	8	12	86	
FMILMLTHF	2075	9	12	86	
FMYVGSVEFQ	6335	11	13	93	
SFSFLLAL	173	9	14	100	0.0041
SFSFLLAL	173	10	14	100	
SMLTDPSHI	2178	9	14	100	
SMDDQAMMKQL	1608	9	11	79	
SMKGSSGQPL	1164	11	12	86	
TWMNSTGF	556	8	11	79	

HCV  $\Lambda$ 24 Motif With Binding Information

Sequence	Position	No. of Amino Acids	Sequence Frequency	Conservancy (%)	A*2401
TWLVGGML	1664	9	1.2	0.6	
TYSTYGKF	1297	0	1.3	9.3	
TYSTYGKF	1297	9	1.2	86	
VFGGLTHI	1566	0	1.3	9.3	
VFGGLTHI	1566	0	1.3	7.9	
VMGSSYGF	2639	0	1.1	1.1	
VMGSSYGF	2639	11	1.3	9.3	
VYLLFRAGPQL	34	8	1.4	100	
WMNRLLAF	1920	10	1.4	100	
YYRGLDVSVI	1422	2			
					53

Table XIX a  
IUCY DLS-Sufer Motif

Core Sequence	Core Freq.	Core Consistency (%)	Exemplary Sequence	Position in Iucy Poly-protein	Iucy Sequence Frequency	Exemplary Sequence Consistency (%)
FGWASWV	12	66	TGFGAYNSKAVDQ	1266	6	36
FGGWAWNST	12	66	GNFGCTTMWNST	550	11	79
FGRKALGIL	12	66	AEGRKALGIL	1730	12	79
FLLALSLCL	12	66	FSFLALSLCLVPA	114	6	86
FPLDGLVRC	11	79	UVPLDGLVRC	2612	11	43
FQVAVMILHAP	12	66	POFQVAVMILHAP	1225	8	79
FRANCTRC	12	66	YDFRANCTRCVK	1182	7	43
FSPFLALL	14	100	GGFSRSPFLALLSLC	171	12	50
FSQPFETI	14	100	TVFSQSFETI	1456	12	86
FTEAMTRHS	14	100	URFTEAMTRHS	2789	7	50
FIPSPOVQ	13	93	WYDFTPSPOVQDID	509	13	93
FITFLPMLST	11	79	POSETFLPMLST	661	9	61
FVKWVAKAF	12	66	IEFVKWVAKAF	1762	3	21
FIQHFLSOT	14	100	LTIQHFLSOT	1570	7	50
IQDQGCTCQ	12	66	DSNDQGCTCQDQ	1454	12	86
IDTTCGFA	12	66	GRMDTTCGFA	120	12	86
IEANLILWRO	12	66	AQLEANLILWRO	2233	7	50
IFLLALSLC	14	100	SFISFLALSLC	173	6	43
FIQKWAQAO	12	66	LFNFGKWAQAO	1013	8	57
IGISGISTD	12	86	STIIGISGISTD	1328	8	57
ISPRNGRS	11	79	CVNQPRNGRS	1903	11	29
ILSPGALVV	13	93	LPAILSPGALVV	1088	11	79
INAYTGPQ	12	66	TEPHWYTTGPQ	2064	8	57
IPLYGARQG	11	79	MCYIYAGARQG	134	10	71
ITTHESEM	12	66	GRNTIVHESEM	2247	10	71
ITSSSSAWS	14	100	LEQITSSSSAWS	2010	11	79
IVFDFLQVN	11	79	AIHDFPFLQVN	2610	11	79
IVAHAYCQL	12	66	QENVIAHAYCQL	1669	6	57
IVDGCSSQG	11	79	GRVLDGCSSQG	1302	10	71
IVLGLSTPQ	14	100	IVYVGLSTPQ	1717	14	100
IVQYQVGA	12	66	IVQYQVGA	1854	10	71
IVATATPPGS	12	66	IYVATATPPGS	1348	9	64
IVCPFETIET	12	66	IVSLOCPFETIET	1466	5	38
IVDQDQIAVAG	12	66	IVVLDQDQIAVAG	1335	12	86
IVETSCSS	13	93	IVYDLEIVETSCSS	2810	13	93
IVFVTSIVW	12	66	SADLEVIVTSIVW	1655	11	79
IVFLFLDADHR	14	100	IVVFLFLDADHR	724	4	29
IVGQWAAQI	12	66	IVNIGQWAAQI	1614	8	57
IVGTYVDO	13	93	IVVIGTYVDO	1329	8	64
IVHYPATRKT	12	66	IVPQHYPATRKT	41	10	71
IVLWVCERM	14	100	IVPQLWVCERM	2615	11	79
IVLGLSAEFL	11	79	IVPFLGLSAEFL	2916	6	43
IVQFSPFLY	12	66	IVTQFSPFLY	1620	11	79
IVQHIVDVO	11	79	IVVQHIVDVO	694	10	71
IVYSPQEI	14	100	IVVYSPQEI	2924	11	79
IVAFASFGN	12	66	IVVIAFASFGN	1921	12	86
IVEANLILWRA	14	100	IVVDEANLILWRA	2232	7	50
IVFCSHOK	14	100	IVVFLCSHOK	1393	14	100
IVTSCSSAV	14	100	IVVLEISCSSSAV	2012	13	93
IVLALLSLCL	12	66	IVVLLALLSLCL	176	5	36
IVLFLLLADAV	14	100	IVVLLFLLLADAV	723	5	29
IVFAGIGGW	12	66	IVVLLFAGIGGW	1609	4	64
IVLLADAVRC	13	93	IVVLLFLLLADAVRC	726	9	71
IVPLAISPG	13	93	IVVLLPLAISPG	1664	10	71

Core Sequence	Core Freq.	Core Consistency (%)	Exemplary Sequence	Position in HCV Poly-protein	Exemplary Sequence Frequency	Consistency (%)
LNQYPLVQ	11	79	FAQDAMQPLVQAK	130	79	79
LNPSVAAATL	14	100	VLVAMPSTAALEQG	1256	100	79
LPALNSFGA	13	93	TTTPALNSFGA	1085	111	79
LPALSTGLI	12	66	TTTPALSTGLI	604	111	79
LPFPLFGAG	13	93	VLVLPDPLFGAG	34	13	93
LIDLAVAVE	11	79	HNGLTILAVAVEPVV	966	4	29
UHKLQPLPPL	12	86	ASCLKLQPLPPLW	2939	7	50
LSAFESAHY	11	79	UCLSAFESAHYSPG	2919	11	79
LSAFSLKAT	11	79	ASOLSAFSLKATCT	2208	7	50
LSNSLRLH	12	86	IVALNSLRLHINAVV	2476	4	20
LSPGALVVA	13	93	PAIISPGALVVA	1089	11	79
LSPLLSST	11	79	RSPLSPLSTTEWQ	664	7	50
LSPLSSTT	11	70	QWLSPLSSTTENWQ	95	11	79
LSTGILIAM	12	65	IVLALSGILIAM	6017	10	71
LTIG-AGLM	12	66	IDLTCG-AGLM	123	12	86
LTHIDANIFL	13	93	FTDIDANIFL	1567	13	93
LTSKLTOPS	13	83	VANLISAKTFSIT	2173	9	64
LYVQDATVCG	12	86	FLVPLVQDATVCGA	1508	9	64
LVQDLAGYCG	11	79	GRVQDLAGYCGAV	1850	9	64
LYGGVLAAL	12	66	TWVWVGGVLAALAY	1664	12	86
LYVFLSVNA	14	100	YVFLVFLSVNAVQ	1254	14	100
LVNLPLVAIL	11	79	TEQDVLNLPLVAIL	1881	10	71
LYVLUQAVI	11	79	DLYLVLVLUQAVI	1134	11	79
LYWAVNCAA	12	66	PDALWVNCAA	1094	11	79
LVVLATIP	12	66	QWALVVLATIPG	1345	11	79
LWATWMLAT	12	60	APWLWATWMLATIF	2069	11	79
LWEDMCTON	12	60	ANLWIDQEMGNTIN	2238	12	86
LYTILQONION	11	79	IMLYLQONION-ET	1627	9	64
WAKRNEWCV	12	66	TRWAKRNEWCV	2569	9	64
WAWDQWAWWV	12	66	GTWAWDQWAWWV	315	12	86
WCGCNTTVE	12	66	NDGCGCNTTVESEN	2213	12	86
WASYIRLYG	11	79	ADLACVYPLVQDQ	131	11	79
WATLDFSLIT	14	100	LTFLMDFSLITAEI	2176	8	57
WAWTRUAFAS	11	79	WAWTRUAFAS	1910	14	100
WAWYDPSGQ	14	100	TERAWYDPSGQ	2783	10	71
WAWYDPSGQ	14	100	AKI AWYDPSGQ	1767	12	86
WAWYDPSGQ	14	100	KWAWYDPSGQ	633	5	36
WAWYDPSGQ	14	100	GAWYDPSGQ	1861	7	50
WAWYDPSGQ	14	100	TTWAWYDPSGQ	1227	6	43
WAWYDPSGQ	14	100	WVWAWYDPSGQ	1437	6	43
WAWYDPSGQ	12	86	PWVWAWYDPSGQ	1589	10	71
WCAVLRH	11	79	WESWFTGLTRH	1899	10	71
WCAVLRH	14	100	WESWFTGLTRH	1563	6	43
WCAVLRH	12	86	WAWGQWAWLAAYC	1665	12	86
WCAVLRH	13	93	CONGCONLUTGSP	2619	13	93
WCAVLRH	12	86	GLPYCQDLFEMESV	1552	6	43
WCAVLRH	11	79	FLAVGCTVWVNGDF	1186	11	79
WCAVLRH	12	86	KANVCFQDFBEGKA	2594	11	79
WCAVLRH	11	79	RSFENFTGCRSPVQ	1211	12	86
WCAVLRH	13	93	WESWFTGLTRH	1563	6	43
WCAVLRH	12	86	WAWGQWAWLAAYC	1665	12	86
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WCAVLRH	12	86	KANVCFQDFBEGKA	2594	11	79
WCAVLRH	12	86	RSFENFTGCRSPVQ	1211	12	86
WCAVLRH	12	86	WESWFTGLTRH	1563	6	43
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WCAVLRH	12	86	GLPYCQDLFEMESV	1552	6	43
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WCAVLRH	12	86	KANVCFQDFBEGKA	2594	11	79
WCAVLRH	12	86	RSFENFTGCRSPVQ	1211	12	86
WCAVLRH	12	86	WESWFTGLTRH	1563	6	43
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WCAVLRH	12	86	GLPYCQDLFEMESV	1552	6	43
WCAVLRH	11	79	FLAVGCTVWVNGDF	1186	11	79
WCAVLRH	12	86	KANVCFQDFBEGKA	2594	11	79
WCAVLRH	12	86	RSFENFTGCRSPVQ	1211	12	86
WCAVLRH	12	86	WESWFTGLTRH	1563	6	43
WCAVLRH	13	93	WAWGQWAWLAAYC	1665	12	86
WCAVLRH	11	79	CONGCONLUTGSP	261		

## HCV DRSuper Motif Binding Data Not Included

Core Sequence	Core Freq.	Core Conservancy (%)	Exemplary Sequence	Position in HCV Polyprotein	Exemplary Sequence Conservancy (%)
VI, AALAAAC YATKIPPG	12 13	86 93	VGGVLAALAAAYCLTT FLVNLATKTPPSGTT	1668 1347	57 64
YLEDGVVAA VLFPSVAT	12 14	86 100	GVRLVEDGQVATGNN KVLVLPNSVAAATLGF	1554 1255	66 100
VLTSMITCP VLTSSGNT	13 11	93 79	DVAVLTSMLTDPFSHI ASGNTTSGCGNTIC	2172 2734	14 10
VLDLAGY VLYGGVLA	12 14	86 100	IGKVVVYVAGYGAG STWVLVGVGVLAAALAA	1849 1663	10 12
VLVLPNSVA VNLNPAILA	12 14	86 86	GKYNVVLNPSVAAATL EDLVNLPAISPGAA	1253 1882	14 14
VPEEDAAAR VSTMPVLG	12 11	86 86	THYPPESDAAARVTO LEVNTSTVVLVGGVIL	1937 1658	11 12
WATDALT WCAALRMT	12 11	79 79	DVNNVATDALTGT WVGVCAALRMT	1436 1898	6 10
WGVYCAAR VLLATTAFF	11 12	79 86	GALVYVCAATLAR AHLVYLATKPPSYY	1895 1346	10 11
WVCFPSV WAGNWLSPR	13 12	93 86	CGPPYCFTRSPVWGG GC3WNGWGLSPRGSR	5056 90	7 5
WHRMMLWTH WGADTAAACG	12 12	86 86	PTLWRLWLNTHFS ITWGDIAACGDI	2870 988	7 6
WGSDPAAFR WAWRJLFA	12 14	86 100	PPSMWGPDFPFRSRIN AVDWMRFLAFAASG	104 1917	10 14
WLLAPITA WTGALTTC	11 12	79 86	SKGIVLALATVIAQ SYTWGALTPTVQ	1025 2456	4 9
WVELTAAET YATSNLPGC	12 13	86 93	GCAYVYELTAFTVNR GMMATCNPCCSFS	1529 161	5 11
YCFPSVW YDAGCAYNE	11 14	79 86	GPVVCFTSPVWGT CECYDAGCAYNE	5077 1523	13 10
YDLCDEC YOLEL1SC	12 13	93 86	GGYDAGCDECEH OPEDLEUTCSSEN	1312 2808	10 10
YGAGVAGL YGFYSPGQ	12 11	86 86	PLSYGKFLDVSQFV GSSYGRSPSFCRVE	1857 2641	11 10
YGFYLGQG3 YKVLVLMPS	11 14	79 100	YSGYKFLDVSQFV AGKIVLVLNPSVAA	1298 1251	10 10
YLAGLSTLP YLGSSGCP	12 12	86 86	GIOVFLGSLTLPGRK PYSTLKGSSGCPUC	1776 1162	11 6
YLTHDPTP YONTVACRA	11 13	79 93	RVYVYLTDPITPLAR LVAVYVAVCAHADAP	1857 1591	13 11
YRGLDVS YRGLAVONE	14 11	100 79	YAYFRGFLDVSMPITS PFLYFLGAVONEVTL	1420 1628	50 7
YRGLAVONE YRGLAVONE	13 13	93 93	NGYFRGRASGNTT GACYSPEELDPAI	1251 2726	9 10
YSEPLDLP YSPCENR	11 11	79 79	UYSPEENRASC UYSPEENRASC	2302 2827	43 57
YUGDCGSV YGYLUPNR	12 11	86 79	SAKYGDCGSVFLY SAKYGDCGSVFLY	3036 3036	8 8

Table XIXb. IICY DR Super Motif With Binding Data

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UCY DR Super Motif With Binding Data

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## HCY DIR Super Motif With Binding Data

Core Sequence	Exemplary Sequence	DR1	DR2w2.1	DR2w2.2	DR3	DR4w4	DR5w5	DR6w6.1	DR6w6.2	DR6w6.3	DR7	DR8	DR9	DR10w10
VLEDGVAYA	DRIVLEDDQVAVYAT	0.0007												-0.0002
VLNPSVAAAT	KVLYLNPSVAAATLOF													
VLYSMATDIP	DYANVTSMLTDPSH													
VLTSGCQT	ASQVITSGCQTLC													
VLVQDLAGY	LGKVLVQDLAGY													
VLVGVGVLA	STVVLVGVGVLA													
VLVLPMSIA	GTKLVLVLPMSIA													
VNLFLPAILS	EDLVNLFLPAILS	1.0000	0.0260	0.0004	0.0300	9.6000	0.0570	0.1400	0.0320	0.6900	0.1700	0.2600	0.0015	1.4000
VPESDAAM	THVNPESDAAMVTO	0.3700				0.0110								
VTSWVLYV	LEWVTSWVLYV	0.0120	0.0078	-0.0003		0.0200		0.0008		0.0016		0.1600	0.0120	
VVATDALMT	DVWVATDALMTGTY	0.0110	0.0110	-0.0003	0.0100	0.0072		-0.0004	0.0140	-0.0003		0.0910	-0.0025	
VVCAAILTR	VVGVVCAAILTR													
VVWVCAW	GALVWVCAW	0.0170												
VVLTATPP	ATLVVLTATPPSV													
VVCFPSV	CCPVCFPSV	0.2700	0.0025	-0.0003		0.0200	0.0005			-0.0001	0.0011	0.2700	0.4300	
VAGWILSPSI	GGGWGWLSPGCSN													
VANWLMH1	PTLVWANWLMH1													
VROADTACQD	ITVHQDITACQD	0.0064				0.0200						0.0190		
VAPDTPHHR	PPSNGPTDPHHR													
VWNRLLFA	AVWNRLLFA	2.2000												
VWRLAPTA	SKRNBLLAPTA	14.0000	0.0730	0.0000	-0.0006	2.0000	0.2500	4.2000	0.0290	-0.0001		0.3900	0.0260	0.0630
WTGALTIP	SYTWGALTIP	0.0250	0.0007	0.0016		0.0560	0.0220	0.0331		-0.0001	0.0130	0.4100	0.0750	
WTETLPACT	CCWVETLPACT													
WTGNLPGC	QVNNNTGNLPGC													
WTGTSPAV	SPVGTGTSPAV													
YDNGCAYYE	CEVYDNGCAYYE													
YDNCDEC	GRAYDNCDEC													
YDLELTSC	DPEVDELTSC	0.0003												
YDAGVAGR	LAQYDAGVAGR	0.0410												
YDOKFSDO	QSSYDOKFSDO	0.4600												
YDPLAZGQ	YSDYKBLDQG	0.0001												
YDSSDGP	ADGDKYLVUNPSVA	0.8400	0.0140	0.0004	0.0045	6.3000	0.1700	0.2700	0.0370	0.5900	0.2800	0.3300	0.2600	
YLTADTPP	PMVNLTRDTPPLR													
YVATVCHARA	LVYAVKLYVCAVHQA													
YVLDVSN	VAYYRGGLDVSMPVTS													
YVLYGAVONE	PLVYLGAVONEVTL													
YVPCRSVN	NCVYTPCRSGVLT													
YSIEPLDLP	GNCYSEPLDLP													
YSPOENIRW	UFSYSEGENINASC													
YVGDGOSV	SAMVNGDGCOSV													
YGTLPLPR														

ICCV 2011 Super-Motif With Binding Data

Table XXb  
HCV\_Dr3Δ Motif Y With Binding Information

Core	Sequence	Exemplary Sequence	DR3	DR1	DR2w2B1	DR2w2B2	DR4w4	DR4w5	DR5w15	DR5w11	DR5w12	DR6w19	DR7	DR8w2	DR9	DRw53
FL00GGCS3	YGRFLADGCCGCGAY															
FL00GPIT1	TVDFSLDPTFIEITT	0.0001														0.0005
LEGEFDPO	NPFLEGEPPGPDPOLSD	-0.0017														
LPCEPEPDV	GSQLPCEPEPDVAVL	-0.0017														0.0230
MANDHMMW	GHEMWADMAMMAMWSP															
MATDPSHIT	LTSMLTOPSHITAE															
MSADLEV/T	MAGMSADLEV/TSTW															
VATDAUWTG	VVVVATDAUWTG	0.0004														-0.0003
VCODHLEFW	GLPVICDHLFWESV	1.1000	0.0046	0.0047	0.0014				0.0006			0.0029	0.0400	0.0029		
VFPDLGWRV	RLNFPDLGWRVCEK	0.0063														
VFTDNNSSPP	RSPVFTDNNSSPPAVP															
VLCECYDAG	DSSVLCCECYDAGCAW	-0.0017														
VLEDGVWYA	GYTMVLEDGVNNTGN															
VLVDILAGY	LGVLVVDILAGYGAQ															
VQFEGGFK	VFCVQFEGGFKPQR															
YDEELTSC	QPEVDQELTSCSSN															
YSEPLDLP	YDCEYSEPLDLPQI															
YVGDLOGSV	SAMVYDLOGSVFLY															
YVFESDAAA	PTIHYVFESDAAAAT															
	19															

Table XXc HCV 3B Motif

Core Sequence	Core Freq.	Core Conservancy (%)	Exemplary Sequence	Position In HCV Poly-protein	Exemplary Sequence Frequency	Exemplary Sequence Conservancy (%)
FQ ISRCCD	14	100	1URCHSRKQICELA	1395	14	100
FSTDTTDFD	11	79	PAFDSYDNTDFPSTV	2667	11	79
LAEGFRKKA	12	86	GHQLAEGFRKKAQG	1728	8	57
UQPHLHAPT	11	79	URQPHLHAPTLL	1616	71	71
VRATRKISE	11	79	FLGIVATRKISEFSO	43	10	71
YLTPLHADV	12	86	SDLYLVTLHADVPPV	1133	11	79
MSNTPROR	11	79		1		

Table XXd IICY3B Motif Binding Data

Core Sequence	Exemplary Sequence*	DRI	DR2w21	DR2w22	DR3	DR4w4	DR4w5	DR5w11	DR5w12	DR6w2	DR6w18	DR7	DR9	DRw53
FDSRKCD	HUFCHSKKICDELA PMYFSTOTRQDSTV GNCZLAEQKQKALG. URUKPTLHSPPLL													
FSTDTRFD	FILQVATRKTQTSRQ	0.0190												
LAEDRDKA	SDLYLYTRHADIVP													
IKPTLHSP														
VRATKTE														
YLVTRHADV														
KSTMRPQR														
7														

TABLE XXI. Population coverage with combined HLA Supertypes

<u>HLA-SUPERTYPES</u>	PHENOTYPIC FREQUENCY					Average
	Caucasian	North American Black	Japanese	Chinese	Hispanic	
<u>a. Individual Supertypes</u>						
A2	45.8	39.0	42.4	45.9	43.0	43.2
A3	37.5	42.1	45.8	52.7	43.1	44.2
B7	38.6	52.7	48.8	35.5	47.1	44.7
A1	47.1	16.1	21.8	14.7	26.3	25.2
A24	23.9	38.9	58.6	40.1	38.3	40.0
B44	43.0	21.2	42.9	39.1	39.0	37.0
B27	28.4	26.1	13.3	13.9	35.3	23.4
B62	12.6	4.8	36.5	25.4	11.1	18.1
B58	10.0	25.1	1.6	9.0	5.9	10.3
<u>b. Combined Supertypes</u>						
A2, A3, B7	83.0	86.1	87.5	88.4	86.3	86.2
A2, A3, B7, A24, B44, A1	99.5	98.1	100.0	99.5	99.4	99.3
A2, A3, B7, A24, B44, A1, B27, B62, B58	99.9	99.6	100.0	99.8	99.9	99.8

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Table XXII HCV ANALOGS

AA	Sequence	Fixed Name.	A1 Motif	A2 Super Motif	A3 Super Motif	A24 Motif	B7 Super Motif	1° Anchor Fixer
9	RVXEKMALY							
9	AVXTRGVAK							
9	EVFXVQPEK							
9	HIFXHSKK							
9	LPGXSFIF							
9	LIFXHSKK							
9	VLAALAAAXL							
10	HIFXHSKK							
10	AAANWTRGEA							
10	YLPRARGPRV							
9	FPGCSFSF							
9	LPVCSFSF							
9	LPGCSFSF							
9	LPGCMFSF							
9	LPFCFSIF							
9	LPGCSFSF							
9	LPGCSFSI							
9	PPVWHGCP							
9	KPTLHGPPTI							
10	APTLWARMII							
9	SPRGSRPSI							
10	LPGRGFLGI							
9	SPCGRVEEI							
9	LPGCSFSI							
9	DPRASHSN							
9	SPGALVGV							
10	TPLLYFLGAI							
9	TISGVLWQV							
9	SISGVLWQV							
9	SLMAFTASV							
9	GLHDCTIMV							
10	KLVALGVNAV							
10	YLPSRGPKL							
10	KLSGLGLNAV							
10	YLPRRGPRL							
10	VFFNLLGKAV							
10	KLVSLGVNAV							
9	CINGYCYTA							
9	CANGVCWTV							

## LUCY ANALOGS

AA	Sequence	Fixed Nomen.	A <sup>1</sup> Molli	A <sup>2</sup> Super Molli	A <sup>3</sup> Super Molli	A <sup>24</sup> Molli	B <sup>7</sup> Super Molli	1 <sup>a</sup> Anchor Fixer
	CVNGVCWAV							

Table XXIII. Immunogenicity of identified supermotif-bearing peptides

Supermotif	Peptide	Sequence	Protein	Position	Human <sup>a</sup>			Immunogenicity			Transgenic mice <sup>b</sup>	
					Barnaba; patients	Barnaba; contacts	Chisari	Page	overall	Frequency	Response	
A2	1073.05	LLFNILGGWV	NS4	1812	1/6	7/17	2/21	0/6	10/50	6/6	6.4 (1.7)	
	1090.18	FILLADARV	NS1/E2	728	2/6	7/17	1/21	0/6	10/50	5/6	9.5 (3.0)	
	1013.02	YLVAYQATV	NS4	1590	1/6	4/17	1/21	0/6	6/50	5/6	8.5 (3.7)	
	1090.22	RLJVFPDLGV	NS5	2578	2/6	5/17	0/21	0/6	7/50	0/6	-	
	1013.1002	DLMGGYPLV	Core	132	2/6	7/17	1/21	1/6	11/50	5/6	8.8 (2.6)	
	24.0073	WMNRLLIAFA	NS4	1920	1/6	3/17	2/21	1/6	7/50	0/6	-	
	24.0075	VLVGGVLAAL	NS4	1666	1/6	6/17	3/21	1/6	11/50	0/6	-	
	1174.08	HMWNFISGI	NS4	1769	3/6	3/17	2/21	0/6	8/50	6/6	6.4 (1.7)	
	1073.06	ILAGYGGAGV	NS4	1851	2/6	3/17	0/21	0/6	5/50	3/6	54.7 (3.3)	
	1073.07	YLLPRRGGPRL	CORE	35	2/6	5/17	7/21	1/6	17/50	4/6	59.1 (7.2)	
A3	24.0071	LLFLLLADA	NS1/E2	726	2/6	9/17	0/21	0/6	11/50	0/6	-	
	1.0119	YLVTRHADV	NS3	1131	6/6	10/17	0/21	1/6	17/50	0/6	-	
	1.0952	KTSERSQPR	CORE	51	2/16	1/4	3/12	0/6	6/38	3/6	23.4 (1.3)	
	1073.11	RLGVVRATRK	CORE	43	4/16	1/4	7/12	1/6	13/38	3/6	42.2 (1.2)	
	1.0955	QLFTFSPPR	ENV	290	1/16	0/4	6/12	1/6	8/38			
B7	1073.13	RMYVGGVEHRR	NS1/E2	632	5/16	1/4	4/12	1/6	11/38	2/6	2.8 (1.1)	
	1.0123	LIFCHSKKK	NS3	1396	6/16	1/4	4/12	2/6	13/38	3/6	4.4 (1.1)	
	1073.10	GVAGALVAFK	NS4	1863	3/16	0/4	6/12	2/6	11/38	6/6	56.5 (1.7)	
	24.0090	VAGALVAFK	NS4	1864	4/16	1/4	6/12	0/4	11/38	1/6	7.1	
	24.0086	TLGFGAYMSK	NS3	1262	6/16	2/12	2/5	10/33				
	1145.12	LPGCSFSIF	CORE	169		2	3/10	5				

**Table XIV. Human and murine MHC-peptide binding assays established using purified MHC molecules and gel filtration chromatography**

A. Class I binding assays						
Species	Antigen	Allele	Cell line	Radiolabeled peptide		Notes
				Steinlin	Source	
Human	A1	A*01:01	Hu. I chain 102-110	YTAVVPLVY		
	A2	A*02:01	HBVc 18-27 F6->Y	FLPSDYFPSV		
	A2	A*02:02	P815 (transfected)	FLPSDYFPSV		
	A2	A*02:03	FUN	FLPSDYFPSV		
	A2	A*02:06	CLA	FLPSDYFPSV		
	A2	A*02:07	721.221 (transfected)	FLPSDYFPSV		
	A3		GM3107	HBVc 18-27 F6->Y		
A11	A11		BVR	non-natural (A3CON1)		
	A24	A*24:02	KAS116	non-natural (A3CON1)		
	A31	A*31:01	SPACH	non-natural (A3CON1)		
	A33	A*33:01	LWAGS	non-natural (A3CON1)		
	A28/68	A*68:01	C1R	HBVc 141-151 T7->Y		
	A28/68	A*68:02	AMAI	HBV pol 646-654 C1->A		
	B7	B*07:02	GM3107	A2 signal seq. 5-13 (L7->Y)		
	B8	B*08:01	Steinlin	IV <sub>BP</sub> 586-593 Y1->F, Q5->R 60s		
	B27	B*27:05	LG2			
	B35	B*35:01	C1R, BVR	non-natural (B35CON2)		
	B35	B*35:02	TISI	non-natural (B35CON2)		
	B35	B*35:03	EHM	non-natural (B35CON2)		
	B44	B*44:03	PITOUT	EF-1 G6->Y		
	B51	B*53:01	KAS116	non-natural (B35CON2)		
	B53	B*53:01	AMAI	non-natural (B35CON2)		
	B54	B*54:01	KT3	non-natural (B35CON2)		
	Cw4	Cw*04:01	C1R	non-natural (C4CON1)		
	Cw6	Cw*06:02	721.221 transfected	non-natural (C6CON1)		
	Cw7	Cw*07:02	721.221 transfected	non-natural (C6CON1)		
Mouse	D <sup>b</sup>		EL4	Adenovirus E1A P7->Y	SGPSNTYPEI	"
	K <sup>b</sup>		EL4	VSV NP S2-59	RGYVFQGL	"
	D <sup>d</sup>		P815	HIV-IIB ENV G4->Y	RGPYRAFTI	"
	K <sup>d</sup>		P815	non-natural (KdCON1)	KFNPMPKTYI	"
	L		P815	HBV <sub>s</sub> 28-39	IPQSLDSYWTSL	"

**Table XIV. Human and murine MHC-peptide binding assays established using purified MHC molecules and gel filtration chromatography**

**B. Class II binding assays**

Species	Antigen	Allele	Cell line	Radio-labeled peptide		Notes
				Source	Sequence	
Human	DR1	DRB1*0101	LG2	HA Y307-319	YPKYVKQNTLKLAT	
DR2	DRB1*1501	L466.1	MBP 88-102Y	VVHFFKAVNTPRIPYY		
DR2	DRB1*1601	L242.5	non-natural (760.16)	YAAFAAAAKTAAAFAA		
DR3	DRB1*0301	MAT	MT 65kD Y3-13	YKTIAFDEEAR		optimal assay pH is 4.5
DR4w4	DRB1*0401	Preiss	non-natural (717.01)	YARFQSQTTLKQKT		
DR4w10	DRB1*0402	YAR	non-natural (717.10)	YARFQRQTTLKA		
DR4w14	DRB1*0404	BIN 40	non-natural (717.01)	YARFQSQTTLKQKT		
DR4w15	DRB1*0405	KT3	non-natural (717.01)	YARFQSQTTLKQKT		
DR7	DRB1*0701	Pitout	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR8	DRB1*0802	OLL	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR8	DRB1*0803	LUY	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR9	DRB1*0901	HID	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR11	DRB1*1101	Sweig	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR12	DRB1*1201	Herluf	unknown eluted peptide	EALIHQLKINPVVLS		
DR13	DRB1*1302	H0301	Tet. tox. 830-843 S->A	QYIKKANSKFIGITE		
DR51	DRB5*0101	Gm3107 or L416.3	Tet. tox. 830-843	QYIKKANSKFIGITE		
DR51	DRB5*0201	L255.1	HA 307-319	PKVVKQNTLKLAT		
DR52	DRB3*0101	MAT	Tet. tox. 1272-1284	NGQIGNDPNRDIL		
DR53	DRB4*0101	L257.6	non-natural (717.01)	YARFQSQTTLKQKT		
DQ3.1	DQA1*0301/DQB1*0301	PF	non-natural (ROIY)	YAHAAHAHAHAHA		no NEM in PI mix
Mouse	IA <sup>b</sup>	DB27.4	non-natural (ROIY)	YAHAAHAHAHAHA		optimal assay pH is 5.5
IA <sup>d</sup>		A20	non-natural (ROIY)	YAHAAHAHAHAHA		
IA <sup>k</sup>		CH-12	HEL 46-61	YNTDGSTDYGLQINSR		optimal assay pH is 5.0
IA <sup>s</sup>		LS102.9	non-natural (ROIY)	YAHAAHAHAHAHA		
IA <sup>u</sup>		91.7	non-natural (ROIY)	YAHAAHAHAHAHA		
IE <sup>d</sup>		A20	Lambda repressor 12-26	YLEDARRKKKAYEKKK		optimal assay pH is 5.0
IE <sup>k</sup>		CH-12	Lambda repressor 12-26	YLEDARRKKKAYEKKK		optimal assay pH is 5.0

**Table XXXV. Monoclonal antibodies used in MHC purification.**

Monoclonal antibody	Specificity
W6/32	HLA-class I
B123.2	HLA-B and C
IVD12	HLA-DQ
LB3.1	HLA-DR
M1/42	H-2 class I
28-14-8S	H-2 D <sup>b</sup> and L <sup>d</sup>
34-5-8S	H-2 D <sup>d</sup>
B8-24-3	H-2 K <sup>b</sup>
SF1-1.1.1	H-2 K <sup>d</sup>
Y-3	H-2 K <sup>b</sup>
10.3.6	H-2 IA <sup>k</sup>
14.4.4	H-2 IE <sup>d</sup> , IE <sup>K</sup>
MKD6	H-2 IA <sup>d</sup>
Y3JP	H-2 IA <sup>b</sup> , IA <sup>s</sup> , IA <sup>u</sup>

Table XXXVI: HCV-derived conserved high algorithm A\*0201-binding peptides

Peptide	Molecule	1st Position	Sequence	Cons.	A2-superype binding capacity (IC50 nM)					
					A*0201	A*0202	A*0203	A*0206	A*6802	A2 XRN
1073.05	NS4	1812	LLFNILGGWV	85	4.2	113	3.2	19	33	5
1090.18	NS1/E2	728	FLLLADARV	92	18	90	149	247	111	5
1013.02	NS4	1590	YLVAYQATV	85	20	39	16	82	33	5
1090.22	NS5	2611	RLIVFPDLGV	79	56	391	10	370	8000	4
1013.1002	CORE	132	DLMGYIPLV	79	80	4778	204	481	12	4
24.0073	NS4	1920	WMNRLIAFA	100	122	130	3.3	1609	400	4
24.0075	NS4	1666	VLVGGVLA	85	185	331	32	308	3077	4
1174.08	NS4	1769	HMWNFISGI	92	15	10750	77	132	7547	3
1073.06	NS4	1851	ILAGYGGAGV	79	116	143	5.0	755	889	3
1073.07	CORE	35	YLLPRRGPRL	92	125	6143	455	416	10256	3
24.0071	NS1/E2	726	LLFLILLADA	100	217	287	455	3364	3077	3
1.0119	LORF	1131	YLVTTRHADV	85	455	2048	3.6	71	3077	3
24.0065	NS4	1891	ILSPGALVV	92	238	10750	27	1028	3077	2
1013.12	NS1/E2	686	ALSTGGLIHL	85	313	7167	45	18500	10256	2
939.14	NS1/E2	696	HLHQNIVDV	85	500	3071	19	1370	10811	2
1090.21	NS5	2918	RLHGLSAFSL	79	179	782	625	18500	12500	1

**Table XXVII: HCV-derived conserved high algorithm A\*03 and/or A\*11 binding peptides**

Peptide	Molecule	1st Position	Sequence	Cons.	A3-supertype binding capacity (IC50 nM)				
					A*03	A*11	A*3101	A*3301	A*6801
1.0952	CORE	51	KTSERSQPR	92	69	94	67	1813	145
1073.11	CORE	43	RLGVRATRK	79	12	207	429	-	3
1.0955	ENV1	290	QLFTFSPRR	79	15	182	621	3766	3
1073.13	NS1/E2	632	RMYVGGEVHR	100	15	300	95	9667	1778
1.0.23	NS3	1396	LIFCHSKKK	100	20	32	2535	24167	333
1073.10	NS4	1863	GVAGALVAFK	85	28	4	3273	26364	118
24.0090	NS4	1864	VAGALVAFK	85	46	7	3750	11600	258
24.0086	NS3	1262	LGFAYMSK	85	136	21	2950	22308	222
1174.16	NS1/E2	557	WMNSTGFTK	79	208	74	12857	690	1429
1073.14	NS3	1261	TLGRGAYMSK	85	136	98	-	22308	8839
1090.23	LORF	1183	AVCTRGVAK	79	423	240	16364	-	2
1090.24	NS5	2596	EVFCVQPEK	85	13750	222	-	-	18
24.0103	NS1/E2	647	AACNWTRGER	85	36667	429	400	5273	4444
1073.16	NS3	1232	HLHAPTGSGK	85	19	2500	-	-	2857
1073.12	NS3	1395	HLIFCHSKKK	100	423	-	20000	-	1
1090.26	NS3	1395	HLIFCHSKK	100	440	10000	-	-	8000

\* A dash indicates IC50nM &gt;30,000

**Table XXVIII: HCV derived conserved B\*0702 binding peptides****A. High conservancy 9- and 10-mer peptides**

Peptide	Molecule	1st Position	Sequence	Consrv.	B7-supertype binding capacity (IC50 nM)			
					B*0702	B*3501	B*51	B*5301
1145.12	Core	169	LPGCSFSIF	92	28	90	100	114
15.00048	E2	681	LPALSTGGL	85	157	-	2.8	6667
15.0234	NS3	1620	KPTLHGPTPL	79	3.9	-	27500	20000
15.0247	NS5	2835	APTLWARMIL	79	6.3	-	5500	-
15.00042	CORE	99	SPRGSSRPSW	79	14	-	11000	-
15.00039	Core	57	QPRGRQQPI	92	24	-	-	-
15.0218	Core	37	LPRRGPRLGV	92	29	-	6111	4000
15.00060	NS5	2615	SPGQRVFEFL	79	46	-	27500	-
15.00043	Core	111	DPRRRSRNL	85	324	-	-	-
15.00063	NS5	2835	APTLWARMI	79	344	-	4583	-
1292.17	NS5	2317	PPVVHGCPL	79	393	-	-	-
15.0239	NS4	1893	SPGALVVGVV	79	423	-	3438	-
15.0235	NS3	1621	TPLLYRLGAV	92	458	-	6875	909

**Table XXVIII: HCV derived conserved B\*0702 binding peptides****B. Additional HCV derived B7 supermotif peptides.**

Peptide	Molecule	1st Position	Sequence	Consv.	B7-superotype binding capacity (IC50 nM)				
					B*0702	B*3501	B*51	B*5301	B*5401
29.0035	NS3	1378	IPFYGKAI	92	458	-	46	-	50
29.0040	Core	37	LPRRGPRL	92	0.85	-	306	-	5000
29.0036	Core	137	IPLVGAPL	79	13	2250	79	-	2857
16.0187	NS1/E2	680	LPCSFITPA	64	423	24000	9167	-	15
29.0039	Core	169	LPGCSFSI	92	500	200	932	620	6250
15.0219	Core	142	APLGGAARAL	71	9.5	-	-	-	12500
29.0031	NS5	2869	APTLWARM	79	13	-	4583	-	4348
15.0231	NS3	1512	RPSGMFDSSV	71	153	-	-	-	1
29.0085	NS5	2474	LPJNALSNSL	57	220	18000	1170	-	11111
29.0037	NS5	2608	KPARLIVF	85	367	-	3235	-	16667
15.0237	NS4	1789	NPAIASLMAF	71	393	9000	5000	-	1
29.0118	NS5	2869	APTLWARMILM	79	423	-	-	-	3030
29.0042	NS4	1720	LPYIEQGM	85	423	-	1375	-	7692

**C. Engineered analogs of B7 supermotif peptides.**

Peptide	Molecule	1st Position	Sequence	Consv.	B7-superotype binding capacity (IC50 nM)				
					B*0702	B*3501	B*51	B*5301	B*5401
1145.12	Core	169	LPGCSFSIF	92	28	90	100	114	6667
1292.24	Core	169	LPGCSFSII	37	4364	5.3	262	1056	3
1145.13	Core	169	FPGCSFSIF	19	1.6	132	3.2	6.7	5

\* A dash indicates IC50 nM &gt;30,000.

**Table XXIX: HCV-derived A1- and A24-motif containing peptides****A. A1-motif peptides**

Peptide	Molecule	Position	Sequence	Conserv.	HLA-A*0101 binding (IC50 nM)
13.0019	NS5	2922	LSAFSLHSY	79	31
1.0509	NS5	2921	GLSAFSLHSY	79	61
1069.62	NS3	1128	CTCGSSDLY	79	68
24.0093	NS5	2129	EVDGVRLHRY	100	167
13.0016	NS3	1241	KSTKVPAAY	85	1923
1.0125	NS3	1525	CYDAGCAWY	79	4032
24.0008	E1	206	DCSNSSIVY	85	16667
24.0094	NS5	2720	TNSKGQNCGY	100	-
24.0096	NS3	1240	GKSTKVPAAY	85	-
24.0100	NS3	1292	TGAPITYSTY	85	-
	NS3	1263	VAATLGFAY	100	
	NS5	2639	VMGSSYGFQY	79	
	NS5	2640	MGSSYGFQY	79	

A dash indicates IC50 nM &gt;25000

**B. A24 -motif peptides**

Peptide	Molecule	Position	Sequence	Conserv.	HLA-A*2402 binding (IC50 nM)
24.0092	NS4	1765	FWAKHMWNF	85	1.7
13.0075	NS4	1778	QYLAGLSTL	100	250
1073.18	NS1/E2	636	MYVGGVEHRL	92	444
13.0074	NS3	1297	TYSTYGKFL	85	522
13.0134	NS5	2647	QYSPGQRVEF	79	667
24.0091	NS4	1772	NFISGIQYL	100	706
13.0131	Core	135	GYIPLVGAPL	79	2105
24.0108	Core	173	SFSIFLLALL	100	2927
13.0132	NS3	1248	AYAAQGYKVL	79	13333
13.0133	NS4	1859	GYGAGVAGAL	85	-
1174.08	NS4	1769	HMWNFISGI	93	
	E1	317	RMAWDMMMNW	85	
	NS1/E2	635	RMYVGGVEHRL	93	
	NS3	1422	YYRGLDVSVI	100	
	NS3	1468	DFSLDPFTFI	100	
	NS3	1608	SWDQMWKCL	79	
	NS3	1664	TWVLVGGVL	85	
	NS4	1732	QFKQKALGL	85	
	NS4	1732	QFKQKALGLL	85	
	NS4	1765	FWAKHMWNFI	85	
	NS4	1919	QWMNRLLIAF	100	
	NS5	2241	LWRQEMGGNI	85	
	NS5	2669	GFSYDTRCF	79	
	NS5	2875	RMILMTHFF	85	

A dash indicates IC50 nM &gt;25000

Table XXX: Immunogenicity of A2-supertype cross-reactive binders

Peptide	Sequence	Protein	Position	Immunogenicity				Transgenic mice <sup>b</sup>	
				Human <sup>a</sup>		Barnaba; Barnaba;		Barnaba; Barnaba;	
		patients	contacts	Chisari	Pape	overall	Frequency	Response	
1073.05	LLFNILGGWV	NS4	1812	1/6	7/17	2/21	0/6	10/50	6/6 6.4 (1.7)
1090.18	FLFLADARV	NS1/E2	728	2/6	7/17	1/21	0/6	10/50	5/6 9.5 (3.0)
1013.02	YLVAYQATV	NS4	1590	1/6	4/17	1/21	0/6	6/50	5/6 8.5 (3.7)
1090.22	RLIVFPDLGV	NS5	2578	2/6	5/17	0/21	0/6	7/50	0/6
1013.1002	DLMGYIPLV	Core	132	2/6	7/17	1/21	1/6	11/50	5/6 8.8 (2.6)
24.0073	WMNRLIAFA	NS4	1920	1/6	3/17	2/21	1/6	7/50	0/6
24.0075	VLVGGVLA	NS4	1666	1/6	6/17	3/21	1/6	11/50	0/6
1174.08	HMWNFISGI	NS4	1769	3/6	3/17	2/21	0/6	8/50	6/6 6.4 (1.7)
1073.06	ILAGYGAGV	NS4	1851	2/6	3/17	0/21	0/6	5/50	3/6 54.7 (3.3)
1073.07	YLIPRRGPRL	CORE	35	2/6	5/17	7/21	1/6	17/50	4/6 59.1 (7.2)
24.0071	LLFLLLADA	NS1/E2	726	2/6	9/17	0/21	0/6	11/50	0/6
1.0119	YLYTRHADV	NS3	1131	6/6	10/17	0/21	1/6	17/50	0/6

a. Data shown represents the number of positive responses over the total number of patients or contacts examined.

b. Frequency represents the number of positive responses over the total number of mice examined. Response indicates the average magnitude (standard deviation) of the response in positive animals, measured in lytic units.

**Table XXXI: Immunogenicity of A3-supertype cross-reactive binders**

Peptide	Sequence	Protein	Position	Immunogenicity				Transgenic mice <sup>b</sup>		
				Human <sup>a</sup>		Barnaba; Barnaba; Chisari		Pepe	overall	Frequency
1.0932	K1SERSQPR	CORE	51	2/16	1/4	3/12	0/6	6/38	3/6	23.4 (1.3)
1073.11	RLGVVRATRK	CORE	43	4/16	1/4	7/12	1/6	13/38	3/6	42.2 (1.2)
1.0935	QLEFTESPRR	ENV	290	1/16	0/4	6/12	1/6	8/38		
1073.13	RMYVGGVEHR	NS1/E2	632	5/16	1/4	4/12	1/6	11/38	2/6	2.8 (1.1)
1.0123	LIFCHSKKK	NS3	1396	6/16	1/4	4/12	2/6	13/38	3/6	4.4 (1.1)
1073.10	GVAGALVAFK	NS4	1863	3/16	0/4	6/12	2/6	11/38	6/6	56.5 (1.7)
24.0090	VAGALVAFK	NS4	1864	4/16	1/4	6/12	0/4	11/38	1/6	7.1
24.0086	TLGFGAYMSK	NS3	1262	6/16	2/12	2/5		10/33		

a. Data shown represents the number of positive responses over the total number of patients or contacts examined.

b. Frequency represents the number of positive responses over the total number of mice examined. Response indicates the average magnitude (standard deviation) of the response in positive animals, measured in lytic units.

Table XXXII. Candidate HCV-derived HTL epitopes

Selection criteria	Peptide	Sequence	Source	Conservancy	
				Total	Core
A. DR-supermotif conserved 15mers	1283.01	GQIVGGVYLLPPIRGPR	HCV Core 28	93	93
	1283.02	VYLLPPIRGPRLGVR	HCV Core 34	93	93
	1283.03	GWLLSPRGSRPSSWGPT	HCV Core 95	79	79
	1283.04	LGKVIDTLLTCGFADL	HCV Core 119	79	86
	1283.05	IDTLLTCGFADLMLGYI	HCV Core 123	86	86
	1283.06	ADLMGYIPLVGAPLG	HCV Core 131	79	79
	1283.07	GVRVLEDGVNYATGN	HCV Core 154	86	86
	1283.08	GVNYATGNLPGCSFS	HCV Core 161	79	86
	1283.09	GCSFSIFLLALLSCL	HCV Core 171	86	100
	1283.10	GHRMIAWDMMMNWSPT	HCV E1 315	86	86
	1283.11	CGPVYCFTPSPVVVG	HCV NS1/E2 506	93	93
	1283.12	VYCFTPSPVVVGTTD	HCV NS1/E2 509	93	93
	1283.13	GNWFGCTWMNSTGFT	HCV NS1/E2 550	79	86
	1283.14	FTTLPALSTGLIHLH	HCV NS1/E2 684	79	86
	1283.17	DLYLVTRHADVIPVVR	HCV NS3 1134	79	79
	1283.18	RAAVCTRGVAKAVADF	HCV NS3 1186	79	79
	1283.20	AQGYKVLVLPNSVAA	HCV NS3 1251	79	100
	1283.21	GYKVLVLPNSVAAATL	HCV NS3 1253	100	100
	1283.22	VLVLPNSVAAATLGF	HCV NS3 1256	100	100
	1283.23	GTVLDQAETAGARLV	HCV NS3 1335	86	86
	1283.24	GARLVVLATATPPGS	HCV NS3 1345	79	86
	1283.25	GRHLIFCHSKKKCDE	HCV NS3 1393	100	100
	1283.27	DSVIDCNTCVTQTVD	HCV NS3 1454	86	86
	1283.28	TVDFSLDPTFTIETT	HCV NS3 1466	79	100
	1283.30	FTGLTHIDAHFLSQ	HCV NS3 1567	93	93
	1283.31	YLVAYQATVCARAQ	HCV NS3 1591	79	93
	1283.32	KPTLHGPTPLYRLG	HCV NS4 1620	79	79
	1283.33	LEVVTSWVLVGGVL	HCV NS4 1658	86	86
	1283.34	TWVLVGGVLAALAY	HCV NS4 1664	86	86
	1283.35	AEQFKQKALGQIOTA	HCV NS4 1730	86	86
	1283.40	PAISPGALVVGVVCA	HCV NS4 1889	79	93
	1283.41	GALVVGVVCAILRR	HCV NS4 1895	79	79
	1283.42	CAAILRRHVGPGEGA	HCV NS4 1903	79	79
	1283.43	AVQWMNRLIAFASRG	HCV NS4 1917	100	100
	1283.44	MNRLIAFASRGNHVS	HCV NS4 1921	86	100
	1283.48	ANLLWRQEMGGNITR	HCV NS5 2238	86	86
	1283.49	RQEMGGGNITRVESEN	HCV NS5 2243	86	86
	1283.52	ARLIVFPDLGVRVCE	HCV NS5 2610	79	79
	1283.53	FPDLGVRVCEKMALY	HCV NS5 2615	79	100
	1283.54	GVRVCEKMALYDVVS	HCV NS5 2619	79	100
	1283.56	QPEYDLELITSCSSN	HCV NS5 2808	79	93
	1283.57	LELITSCSSNVSVAH	HCV NS5 2813	79	100
	1283.58	PTLWARMILMTHFFS	HCV NS5 2870	79	86
	1283.59	LHGLSAFSLHSYSPG	HCV NS5 2919	79	79
	1283.60	AFSLHSYSPGEINRV	HCV NS5 2924	79	79

Table XXXII. Candidate HCV-derived HTL epitopes

Selection criteria	Peptide	Sequence	Source	Conservancy	
				Total	Core
B. High algorithm conserved core	1283.15	VVLLFLLLADARVCS	HCV NS1/E2 724	29	100
	1283.16	SKGWRLLAPITAYAQ	HCV NS3 1025	29	79
	1283.19	PQTQVVAHLHAPTGS	HCV NS3 1225	43	85
	1283.26	DVVVVATDALMTGYT	HCV NS3 1436	43	79
	1283.29	WESVFTGLTHIDAHF	HCV NS3 1563	43	92
	1283.45	LTSMLTDPSSHITAET	HCV NS5 2176	57	100
	1283.46	ASQLSAPSLSKATCTT	HCV NS5 2208	50	79
	1283.47	DADLIEANLLWRQEM	HCV NS5 2232	50	85
	1283.50	SYTWIGALITPCAAE	HCV NS5 2456	64	79
	1283.51	TTIMAKNEVFCVQPE	HCV NS5 2589	64	85
	1283.55	GSSYGFQYSPGQRVE	HCV NS5 2641	71	79
	1283.61	ASCLRKLGVPPLRVW	HCV NS5 2939	50	85
C. Collaborator	F098.03	AAYAAQGYKVVLNPSVAAT	HCV NS3 1242-1261	71	100
	F098.04	GYKVLVLPNSVAATLGFAY	HCV NS3 1248-1267	100	
	F098.05	GYKVLVLPNSVAAT	HCV NS3 1248-1261	100	
	F134.01	RRPQDVVKFPGGGQIVGGVY	HCV Core 17-35	86	
	F134.02	DVKFPGGGQIVGGVYLLPRR	HCV Core 21-40	86	
	F134.03	GYKVLVLPNSVAATLGFAY	HCV NS3 1253-1272	100	
	F134.04	TLHGPPTLLYRLGAVQNEIT	HCV NS4 1622-1641		79
	F134.05	NFISGIQYLAGLSTLPGNPA	HCV NS4 1772-1791	100	
	F134.06	LLFNIILGGWVAAQLAAPGAA	HCV NS4 1812-1831		86
	F134.07	GPGEGAVQWMNRLLIAFASRG	HCV NS4 1912-1931	86	100
	F134.08	GEGAVQWMNRLLIAFASRGNHV	HCV NS4 1914-1934	100	
	Pape 21	AIPLEVIKGGRHLIFCHSKR	HCV NS3 1379-1398	21	100
	Pape 22	GRHLIFCHSKRKCDDELATKL	HCV NS3 1388-1407		100
	Pape 29	SVIDCNCVQTVDSDLPT	HCV NS3 1450-1469	86	
D. DR3 motif	35.0102	GVRVLEDGVNYATGN	HCV 154	86	86
	35.0103	SAMYVGDLCGSVFLV	HCV 273	57	86
	35.0104	GHRMAWDMMMNWSPT	HCV 315	86	86
	35.0105	SDLYLVTRHADVIPV	HCV 1133	79	86
	35.0106	VVVVATDALMTGYTG	HCV 1437	42	86
	35.0107	TVDFSLDPTFTIETT	HCV 1466	79	100
	35.0108	DSSVLCECYDAGCAW	HCV 1518	71	93
	35.0109	GLPVCQDHLEFWESV	HCV 1552	42	86
	35.0110	GMQLAEQFKQKALGL	HCV 1726	57	86
	35.0111	PTHYVPESDAAARVT	HCV 1936	86	86
	35.0112	GSQLPCEPEPDVAVL	HCV 2162	64	86
	35.0113	LTSMLTDPSSHITAET	HCV 2176	57	100
	35.0114	MPPLEGEPGDPLSD	HCV 2401	79	100
	35.0115	QPEYDLELITSCSSN	HCV 2808	79	93
	1283.25	GRHLIFCHSKKKCDE	HCV NS3 1393-1407		

Table XXXIII. HLA-DR screening panels

Screening Panel	Antigen	Alleles	Representative Assay			Phenotypic Frequencies				Avg.
			Allele	Alias	Cauc.	Blk.	Jpn.	Chn.	Hisp.	
Primary	DR1	DRB1*0101-03	DRB1*0101	(DR1)	18.5	8.4	10.7	4.5	10.1	10.4
	DR4	DRB1*0401-12	DRB1*0401	(DR4w4)	23.6	6.1	40.4	21.9	29.8	24.4
	DR7	DRB1*0701-02	DRB1*0701	(DR7)	26.2	11.1	1.0	15.0	16.6	14.0
	Panel total				59.6	24.5	49.3	38.7	51.1	44.6
Secondary	DR2	DRB1*1501-03	DRB1*1501	(DR2w2 β1)	19.9	14.8	30.9	22.0	15.0	20.5
	DR2	DRB5*0101	DRB5*0101	(DR2w2 β2)	-	-	-	-	-	-
	DR9	DRB1*09011,09012	DRB1*0901	(DR9)	3.6	4.7	24.5	19.9	6.7	11.9
	DR13	DRB1*1301-06	DRB1*1302	(DR6w19)	21.7	16.5	14.6	12.2	10.5	15.1
Tertiary	DR4	DRB1*0405	DRB1*0405	(DR4w15)	-	-	-	-	-	-
	DR8	DRB1*0801-5	DRB1*0802	(DR8w2)	5.5	10.9	25.0	10.7	23.3	15.1
	DR11	DRB1*1101-05	DRB1*1101	(DR5w11)	17.0	18.0	4.9	19.4	8.1	15.5
	Panel total				42.0	33.9	61.0	48.9	30.5	43.2
Quaternary	DR3	DRB1*0301-2	DRB1*0301	(DR3w17)	17.7	19.5	0.4	7.3	14.4	11.9
	DR12	DRB1*1201-02	DRB1*1201	(DR5w12)	2.8	5.5	13.1	17.6	5.7	8.9
	Panel total				20.2	24.4	13.5	24.2	19.7	20.4

**Table XXXIV. HLA-DR binding capacity of target derived peptides: DR-supermotif and algorithm positive peptides.**

Peptide	Sequence	Source	Binding capacity (IC50 nM)									DR alleles bound	
			DR1	DR2w2B1	DR2w2B2	DR4w4	DR4w15	DR5w11	DR6w19	DR7	DR8w2	DR9	
1283.21	AAYAAQGYKVLVLNPSVAATLGFAY	HCV NS3 1242-1267											
	GYKVLVLNPSVAATL	HCV NS3 1253	4.5	350	5.2	567	143	5.1	89	288	54	175	9
1283.20	AQGYKVLVLNPSVAA	HCV NS3 1251	6.0	650	7.9	224	74	5.9	833	175	375	298	9
F98.03	AAYAAQGYKVLVLNPSVAAAT	HCV NS3 1242	2.9	48	483	18	124	103	11	96	60	240	9
F98.05	GYKVLVLNPSVAAAT	HCV NS3 1248-1261	1.4	39	365	7.8	141	75	3.5	126	21	266	9
F98.04	GYKVLVLNPSVAATLGFAY	HCV NS3 1248-1267	3.5	42	854	97	1500	240	4.1	23	80	20	8
	GEGAYQWMNRLIAFASRGNHVS	HCV NS4 1914+1935											
1283.44	MNRLIAFASRGNHVS	HCV NS4 1921	66	4.8	159	322	585	45	73	227	102	313	147
F134.08	GEGAVQWMNRLIAFASRGNHV	HCV NS4 1914	3.2	182	361	345				221	158	16818	6
1283.16	SKGWRLLAPITAYAQ	HCV NS3 1025	0.36	125	23	24	152	4.8			962	54	1190
1283.55	GSSYGFQYSPGQRVE	HCV NS5 2641	11	11	667	417	745	20000	19	156	68	571	7
1283.61	ASCLRKLGVPPLRWW	HCV NS5 2939	5.0	16	217	6250	78	645	2500	862	671	862	-
F134.05	NFISGIGQYLAGLSTLPGNPA	HCV NS4 1772	10	606	84	29				70	441		6

Shading indicates IC50 > 1  $\mu$ M.

A dash (-) indicates IC50 >20  $\mu$ M.

**Table XXXV. HLA-DR binding capacity of 3 DR3 motif-containing peptides**

Peptide	Sequence	Source	DR3 binding (IC50 nM)
35.0106	VVVVATDALMGTGTYG	HCV 1437	427
35.0107	TVDFSLDPFTFTIETT	HCV 1466	235
1283.25	GRHLIFCHSKKKCDE	HCV NS3 1393	ND

Table XXXVIIa: HCV-derived CTL epitope candidates

Peptide	Molecule	1st Position	Sequence	Consy.	Selection criteria
1073.05	NS4	1812	LLFNILGGWV	85	A2-superype
1090.18	NS1/E2	728	FLLADARV	92	A2-superype
1013.02	NS4	1590	YLVAYQATV	85	A2-superype
1090.22	NS5	2611	RLIVFPDLGV	79	A2-superype
1013.1002	CORE	132	DLMGYIPLV	79	A2-superype
24.0073	NS4	1920	WMNRLLIAFA	100	A2-superype
24.0075	NS4	1666	VLVGGVLA	85	A2-superype
1174.08	NS4	1769	HMWNFISGI	92	A2-superype
1073.06	NS4	1851	ILAGYGAGV	79	A2-superype
1073.07	CORE	35	YLLPQRGPRL	92	A2-superype
24.0071	NS1/E2	726	LLFLLLADA	100	A2-superype
1.0119	LORF	1131	VLVTRHADV	85	A2-superype
1.0952	CORE	51	KTSERSQPR	92	A3-superype
1073.11	CORE	43	RLGVRATRK	79	A3-superype
1.0955	ENV 1	290	QLFTFSPPR	79	A3-superype
1073.13	NS1/E2	632	RMYVGGVIEHR	100	A3-superype
1.0123	NS3	1396	LIFCHSKKK	100	A3-superype
1073.10	NS4	1863	GVAGALVAFK	85	A3-superype
24.0090	NS4	1864	VAGALVAFK	85	A3-superype
24.0086	NS3	1262	TLGFGAYMSK	85	A3-superype
F104.01	NS5	3003	VGIYLLPNR	79	A31
1145.12	Core	169	LPGCSFSIF	92	B7-superype
29.0035	NS3	1378	IPFYGKAI	92	B7-superype
13.0019	NS5	2922	LSAFSLHSY	79	A1
1069.62	NS3	1128	CTCGSSDLY	79	A1
24.0092	NS4	1765	FWAKHMWNF	85	A24

**Table XXXVIIb: HCV-derived HTL epitope candidates**

Region	Peptide	Motif <sup>1</sup>	Sequence
HCV NS3 1025-1039	I283.16	DR	SKGWRILLAPITAYAQ`
HCV NS3 1242-1267	F98.03	DR	AYAAAGQGYKVVLNPSVAAT`
HCV NS3 1393-1407	I283.25	DR3	GRHLIFCHSKKKCDE`
HCV NS3 1437-1451	35.0106	DR3	VVVVATDALMTGYTG`
HCV NS3 1466-1480	35.0107	DR3	TVDFSLIDPTFTIETT`
HCV NS4 1772-1790	F134.05	DR	NFISGIQYLAGLSTLPGNPA`
HCV NS4 1914-1935	F134.08	DR	GEGAVQWMNRQIAFASRGNHV`
HCV NS5 2641-2655	I283.55	DR	GSSYGFQYSPGQRV`
HCV NS5 2939-2953	I283.61	DR	ASCLRKLGVPPRLRVW`

1. Peptides identified on the basis of either the DR P1-P6 supermotif or by use of the DR1-4-7 algorithms are indicated by 'DR'. Peptides identified using the DR3 motif are indicated by 'DR3'.

**Table XXXVII. Estimated population coverage by a panel of HCV derived HTL epitopes**

Antigen	Alleles	Representative assay	No. of epitopes <sup>2</sup>	Population coverage (phenotypic frequency)					
				Cauc.	Blk.	Jpn.	Chn.	Hisp.	Avg.
DR1	DRB1*0101-03	DR1	6	18.5	8.4	10.7	4.5	10.1	10.4
DR2	DRB1*1501-03	DR2w2 $\beta$ 1	3	19.9	14.8	30.9	22.0	15.0	20.5
DR2	DRB5*0101	DR2w2 $\beta$ 2	6	-	-	-	-	-	-
DR3	DRB1*0301-2	DR3	2	17.7	19.5	0.40	7.3	14.4	11.9
DR4	DRB1*0401-12	DR4w4	5	23.6	6.1	40.4	21.9	29.8	24.4
DR4	DRB1*0401-12	DR4w15	3	-	-	-	-	-	-
DR7	DRB1*0701-02	DR7	5	26.2	11.1	1.0	15.0	16.6	14.0
DR8	DRB1*0801-5	DR8w2	5	5.5	10.9	25.0	10.7	23.3	15.1
DR9	DRB1*09011,09012	DR9	3	3.6	4.7	24.5	19.9	6.7	11.9
DR11	DRB1*1101-05	DR5w11	5	17.0	18.0	4.9	19.4	18.1	15.5
DR13	DRB1*1301-06	DR6w19	2	21.7	16.5	14.6	12.2	10.5	15.1
<b>Total</b>			<b>98.5</b>	<b>95.1</b>	<b>97.1</b>	<b>91.3</b>	<b>94.3</b>	<b>95.1</b>	

1. Total population coverage has been adjusted to account for the presence of DRX in many ethnic populations. It has been assumed that the range of specificities represented by DRX alleles will mirror those of previously characterized HLA-DR alleles. The proportion of DRX incorporated under each motif is representative of the motif in the remainder of the population. Total coverage has not been adjusted to account for unknown gene types.

2. Number of epitopes represents a minimal estimate, considering only the epitopes shown in Table 6. Additional alleles possibly bound by nested epitopes have not been accounted.

TABLE Ia

SUPERMOTIFS	POSITION	POSITION	POSITION
	2 (Primary Anchor)	3 (Primary Anchor)	C Terminus (Primary Anchor)
A1	<b>T, I, L, V, M, S</b>		<b>F, W, Y</b>
A2	<b>V, Q, A, T</b>		<b>I, V, L, M, A, T</b>
A3	<b>V, S, M, A, T, L, I</b>		<b>R, K</b>
A24	<b>Y, F, W, I, V, L, M, T</b>		<b>F, I, Y, W, L, M</b>
B7	<b>P</b>		<b>V, I, L, F, M, W, Y, A</b>
B27	<b>R, H, K</b>		<b>F, Y, L, W, M, I, V, A</b>
B58	<b>A, T, S</b>		<b>F, W, Y, L, I, V, M, A</b>
B62	<b>Q, L, I, V, M, P</b>		<b>F, W, Y, M, I, V, L, A</b>
<hr/>			
MOTIFS			
A1	<b>T, S, M</b>		<b>Y</b>
A1		<b>D, E, A, S</b>	<b>Y</b>
A2.1	<b>V, Q, A, T*</b>		<b>V, L, I, M, A, T</b>
A3.2	<b>L, M, V, I, S, A, T, F, C, G, D</b>		<b>K, Y, R, H, F, A</b>
A11	<b>V, T, M, L, I, S, A, G, N, C, D, F</b>		<b>K, R, H, Y</b>
A24	<b>Y, F, W</b>		<b>F, L, I, W</b>

\*If 2 is V, or Q, the C-term is not L

Bolded residues are preferred, italicized residues are less preferred: A peptide is considered motif-bearing if it has primary anchors at each primary anchor position for a motif or supermotif as specified in the above table.

WHAT IS CLAIMED IS

1. A composition comprising a prepared hepatitis C virus (HCV) epitope consisting of an amino acid sequence selected from the group consisting of :  
FLLADARV, YLVAYQATV, RLIVFPDLGV,  
DLMGYIPLV, WMNRLIAFA, VLVGGVLAA,  
HMWNFISGI, ILAGYGAGV, YLLPRRGPRL,  
LLFLLLADA, YLVTRHADV, KTSERSQPR,  
RLGVRATRK, QLFTFSPRR, RMYVGGVEHR,  
LIFCHSKKK, GVAGALVAFK, VAGALVAFK,  
TLGFGAYMSK, LPGCSFSIF, LSAFSLHSY,  
CTCGSSDLY, FWAKHMWNF, SKGWRLLAPITAYAQ,  
AYAAQGYKVLVLNPSVAAT, GRHLIFCHSKKCDE, VVVVATDALMTGYTG,  
TVDFSLDPTFTIETT, NFISGIQYLAGLSTLPGNPA,  
GEGAVQWMNRLIAFASRGNHV, GSSYGFQYSPGQRVE, ASCLRKLGVPPRLRVW,  
and LTCGFADLMGY.
2. The composition of claim 1, further comprising two epitopes selected from the group in claim 1.
3. The composition of claim 2, further comprising three epitopes selected from the group in claim 1.
4. The composition of claim 1, wherein the composition further comprises a CTL epitope selected from the group consisting of LTDPHITA, LADGGCSGGAY, RMILMTHFF, VMGSSYGF, FWAKHMWNFI, LLFNILGGWV, IPFYGKAI, and VGIYLLPNR.
5. The composition of claim 1, wherein the composition further comprises an HTL epitope.
6. The composition of claim 5, wherein the HTL epitope is a pan DR binding molecule.

7. The composition of claim 1, wherein the epitope is on or within a liposome.

8. The composition of claim 1, wherein the peptide is joined to a lipid.

9. The composition of claim 1, wherein the epitope is bound to an HLA heavy chain,  $\beta$ 2-microglobulin, and strepavidin complex, whereby a tetramer is formed.

10. The composition of claim 1, wherein the epitope is bound to an HLA molecule on an antigen presenting cell.

11. The composition of claim 10, wherein the antigen presenting cell is a dendritic cell.

12. The composition of claim 1, the composition further comprising a pharmaceutical excipient.

13. The composition of claim 1, further wherein the epitope is in a unit dose form.

14. A composition comprising a prepared peptide of less than 250 amino acid residues comprising at least two hepatitis C virus (HCV) peptide epitopes selected from the group consisting of:

FLLLADARV,	YLVAYQATV,	RLIVFPDLGV,
DLMGYIPLV,	WMNRLIAFA,	VLVGGVLAA,
HMWNFISGI,	ILAGYGAGV,	YLLPRRGPRL,
LLFLLLADA,	YLVTRHADV,	KTSERSQPR,
RLGVRATRK,	QLFTFSPRR,	RYMYVGGVEHR,
LIFCHSKKK,	GVAGALVAFK,	VAGALVAFK,
TLGFGAYMSK,	LPGCSFSIF,	LSAFSLHSY,
CTCGSSDLY,	FWAKHMWNF,	SKGWRLLAPITAYAQ,

AAYAAQGYKVLVLNPSVAAT, GRHLIFCHSKKCDE, VVVVATDALMTGYTG,  
TVDFSLDPTFTIETT, NFISGIQYLAGLSTLPGNPA,  
GEGAVQWMNRLIAFASRGNHV, GSSYGFQYSPGQRVE, ASCLRKLGVPPLRVW,  
and LTCGFADLMGY.

15. The composition of claim 14, wherein at least two epitopes are linked via a spacer.

16. The composition of claim 14, further comprising a third epitope.

17. The composition of claim 16, wherein the third epitope is selected from the group consisting of LTDPHITA, LADGGCSGGAY, RMILMTHFF, VMGSSYGF, FWAKHMWNFI, LLFNILGGWV, IPFYGKAI, and VGIYLLPNR.

18. The composition of claim 16, further comprising a third epitope that is an HTL epitope.

19. The composition of claim 18, wherein the HTL epitope is a panDR binding molecule.

20. The composition of claim 14, wherein the peptide is on or within a liposome.

21. The composition of claim 14, wherein the peptide is joined to a lipid.

22. The composition of claim 14, wherein the peptide further comprises at least three of the epitopes in the group of claim 14.

23. The composition of claim 14, wherein the peptide further comprises at least four of the epitopes in the group of claim 14.

24. The composition of claim 14, wherein the peptide further comprises at least five of the epitopes in the group of claim 14.

25. The composition of claim 14, wherein the peptide further comprises at least six of the epitopes in the group of claim 14.

26. The composition of claim 14, the composition further comprising a pharmaceutical excipient.

27. The composition of claim 14, further wherein the epitope is in a unit dose form.

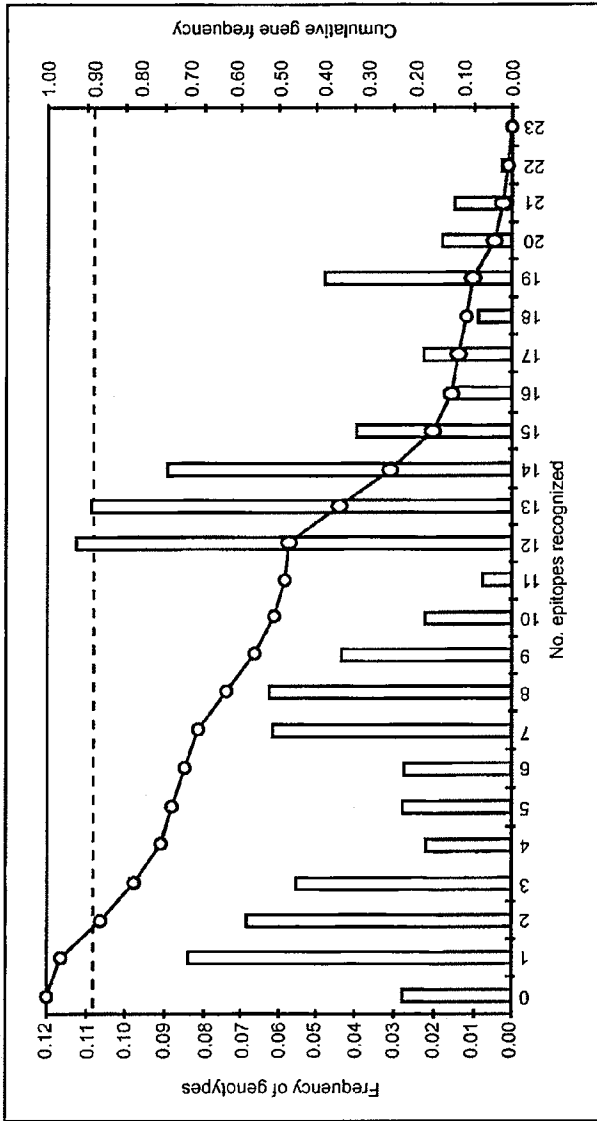
28. A composition comprising at least six prepared HCV epitopes each consisting of an amino acid sequence selected from the group consisting of:

FLLLADARV,	YLVAYQATV,	RLIVFPDLGV,
DLMGYIPLV,	WMNRLIAFA,	VLVGGVLAA,
HMWNFISGI,	ILAGYGAGV,	YLLPRRGPRL,
LLFLLLADA,	YLVTRHADV,	KTSERSQPR,
RLGVRATRK,	QLFTFSPRR,	RMYVGGVEHR,
LIFCHSKKK,	GVAGALVAFK,	VAGALVAFK,
TLGFGAYMSK,	LPGCSFSIF,	LSAFSLHSY,
CTCGSSDLY,	FWAKHMWNF,	SKGWRLLAPITAYAQ,
AAYAAQGYKVLVLNPSVAAT,	GRHLIFCHSKKKCDE,	VVVVATDALMTGYTG,
TVDFSLDPTFTIETT,	NFISGIQYLAGLSTLPGNPA,	
GEGAVQWMNRLIAFASRGNHV,	GSSYGFQYSPGQRVE,	ASCLRKLGVPPLRVW,
and LTCGFADLMGY.		

29. The composition of claim 28, further comprising at least one epitope selected from the group consisting of LTDPHITA, LADGGCSGGAY, RMILMTHFF, VMGSSYGF, FWAKHMWNFI, LLFNILGGWV, IPFYGKAI, and VGIYLLPNR.

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Monte Carlo population coverage analysis for  
HCV candidate epitopes



Plot of total frequency of genotypes as a function of the number of HCV candidate epitopes bound by HLA-A and B alleles, in an average population. Genotype values were derived by averaging the gene frequencies in Caucasian, North American, Black, Japanese, Chinese, and Hispanic populations. Also shown is the cumulative frequency of genotypes.

Using currently available HLA typing data, a residual fraction (about 15%) of the genes, in an average population, are unspecified. To arrive at 100% accounting of genes, a fraction of the residual has been added for each HLA population cluster in proportion to the relative frequency of the cluster within the HLA specified population. One peptide, 24.0086, was not incorporated into the present analysis.

FIG. 1

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## HVC Minigene

CTL Epitopes										Core 132
Kozak	SigSeq	Core 43	NS4 1590	NS3 1128	NS5 2611	Core 169	NS1/E2 632	NS4 1765	NS4 1863	Core 132
A3		1073.11	1013.02	1069.62	1090.02	1145.12	1073.13	24.0092	1073.10	1013.10
		A2	A1	A2	B7	A3	A24	A3	A2	A2
HTL Epitopes										
NS3 1253	NS4 1921	1437	NS5 2641	1466						
1283.21	1283.44	35.0106	1283.55	35.0107						
DR	DR	DR3	DR	DR3						

FIG. 2

**INTERNATIONAL SEARCH REPORT**

International application No.  
PCT/US00/19774

**A. CLASSIFICATION OF SUBJECT MATTER**

IPC(7) : A61K 38/00, 38/04, 38/08, 38/10, 39/29, 39/295

US CL : 514/2,12,13,14,15, 885; 424/185.1, 189.1

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 514/2,12,13,14,15, 885; 424/185.1, 189.1

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

MEDLINE, BIOSIS, EMBASE, DERWENT WPI, WEST 2.0, search terms: author names, hcv, peptid?, HLA, htl, ctl,

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim N
Y	WENTWORTH et al. Differences and similarities in the A2.1-restricted cytotoxic T cell repertoire in humans and human leukocyte antigen-transgenic mice. Eur. J. Immunol. 1996. Vol 26. pages 97-101, see entire document.	1-29
Y	US 5,736,142 A (SETTE et al.) 07 April 1998, see entire document.	1-29

Further documents are listed in the continuation of Box C.  See patent family annex.

• Special categories of cited documents:	*T*	later document published after the international filing date or prior date and not in conflict with the application but cited to understand the principle or theory underlying the invention
*A* document defining the general state of the art which is not considered to be of particular relevance	*X*	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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*O* document referring to an oral disclosure, use, exhibition or other means		
*P* document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search

19 SEPTEMBER 2000

Date of mailing of the international search report

17 OCT 2000

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